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(54) Title: CANDIDA TOPOISOMERASEIGENE

(57) Abstract

The present invention provides an isolated and purified polynucleotide that encodes Candida albicans type I topoisomerase. Methods of making recombinant C. albicans topoisomerase I using those polynucleotides and host cells transformed with those polynucleotides are also provided. The present invention also provides a method for identifying compounds which inhibit the growth of fungal cells using the polynucleotide.

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CANDIDA TOPOISOMERASE I GENE

This application is a Continuation-in-Part Application of and claims the benefit of U.S. Provisional Application 60/005,989, filed October 27, 1995 and also claims the benefit of U.S. Application, filed October 25, 1996, entitled Candida Topoisomerase I Gene to inventors J.M. Fostel. A. Taylor, K.M. Giles, T.P. McGonigal and A. V. Sarthy.

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Background of the Invention

Candida albicans (C. albicans) is an opportunistic pathogen which causes systemic candidiasis in immunocompromised patients. The patient pool at risk includes cancer patients undergoing chemotherapy, organ transplant recipients, patients infected with HIV, and other patients without a healthy immune response. This population is increasing in number annually, making infections by Candida spp. and other fungi an increasingly important clinical challenge (recently reviewed by Kerridge, 1995). It has been estimated that the majority of mycotic infections occurring in cancer patients are caused by Candida albicans and other species of Candida (Kerridge, 1995; Abi-Said and Anaissie, 1995), and that the excess mortality that can be attributed to candidemia is 38% (Wey et al., 1988).

Current therapeutic agents available to combat Candida infections include amphotericin B, a polyene, and the family of azole compounds. Both classes of compounds inhibit fungal growth either through a direct or indirect interaction with an integral membrane component, ergosterol. Despite their effectiveness, however, neither compound completely satisfies therapeutic requirements. Amphotericin B, for example, is associated with adverse reactions in the host (Kerridge, 1995; Rex, et al., 1994). The triazoles, while safer, may have reduced efficacy against a growing population of resistant Candida species. (Kerridge, 1995; DeMuri and Hostetter, 1995; Law et al., 1994). Thus there is an unmet need for safe antifungal agents which also can be used against azole-resistant isolates of Candida spp.

It has been proposed that agents which adversely effect topoisomerases and in particular, type I topoisomerases, may be effective as antifungal agents (Figitt, 1989; Fostel et al., 1992; Shen et al., 1992). DNA topoisomerases are enzymes which modulate the topological structure of DNA and are classified into two classes based primarily on their mode of cleaving DNA. Type I DNA topoisomerases introduce a transient break in one strand of DNA, pass another strand through the nick and change the linking number (i.e. the number of times the DNA backbone strands intertwine) by one unit. Type II topoisomerases introduce concerted breaks in both DNA strands, pass another double-stranded DNA segment through the gap and change the linking number by two. This difference in

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mechanism allows type II topoisomerases to catenate and decatenate covalently closed double-stranded DNA rings (Hsieh and Brutlag, 1980; Kreuzer and Cozzarelli, 1980), and also to knot and unknot covalently closed double-stranded circles (Liu et al. 1980; Hsieh, 1983). The type I topoisomerases, on the other hand, cannot catenate or decatenate double-stranded DNA unless a nick or gap is present in one of the circles (Tse and Wang 1980; Brown and Cozzarelli 1981).

In eukaryotic cells, topoisomerases participate in cellular processes which are associated with the separation of complementary DNA strands (i.e. replication, transcription, recombination and repair). Topoisomerases are also required for the segregation of daughter chromosomes during cytokinesis. Despite the important function these enzymes perform, type I topoisomerases are considered to be non-essential enzymes in that disruption or inhibition of their catalytic activity does not lead to cell death. (It has been proposed that functional topoisomerase II activity may replace or substitute for lost topoisomerase I activity). Thus, selective inhibition of the topoisomerase I catalytic activity of a particular fungal pathogen would not necessarily cure a host of that pathogen. Although type II topoisomerases could prove to be effective antifungal targets, to date, no compounds have been identified which selectively target the cleavage complex formed by the topoisomerase II enzymes of fungal origin.

It has been found however that type I topoisomerases can be adversely affected via a non-catalytic pathway, presumably by "stabilizing" a "cleavage complex" formed between a type I topoisomerase and DNA. In a normally replicating cell, a cleavage complex is characterized by a transient association between the topoisomerase I and DNA during which time the enzyme performs various catalytic functions (such as binding to and nicking a particular region of the DNA, passing a DNA segment through or around the break to change the linking number and ultimately resealing (i.e. repairing) the nick). Stabilizing agents are those chemical compounds which promote cell death through some manner of interaction with a topoisomerase I/DNA cleavage complex, although the nature of this interaction and the mechanism by which the agent induces cell death is unknown. Nitiss et al., for example, have demonstrated the fungicidal effect of camptothecin (which stabilizes the topoisomerase I cleavage complex of Saccharomyces cerevisiae (Nitiss and Wang, 1988) and etoposide and amsacrine (which stabilize the topoisomerase II cleavage complex of Saccharomyces cerevisiae (Nitiss, 1992). Eng et al. have also demonstrated this effect in strains of S. cerevisiae and Schizosacheromyces pombe (Eng, et al., 1988). Thus, it is reasonable to expect a novel class of stabilizing agents to have a similar effect on topoisomerase/DNA cleavage complexes of pathogenic fungi.

In order to minimize unwelcome effects on host tissue, it is important to determine if the fungal topoisomerases have sufficient biochemical differences from the human enzymes WO 97/15676 PCT/US96/17291

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to allow differential targeting by chemical agents. Fostel et al. (1992) isolated a type I topoisomerase from Candida albicans and characterized its response to known topoisomerase agents. The enzyme, Candida topo I, exhibited a differential response (relative to human topo I) to camptothecin and the aminocatechol A-3253 in being ten times less sensitive to camptothecin than human topo I and ten times more sensitive to A-3253 (Fostel et al., 1992; Fostel and Montgomery, 1995). Others have also demonstrated a differential response by type I and type II topoisomerases to different agents (Figgitt et al., 1989; Dykstra et al., 1994).

Efforts to characterize biochemical differences between fungal topoisomerases and host topoisomerases have, however been hampered by the lack of easy and cost-effective methods to assay for such differences. For example, one common assay technique for measuring the effects of stabilizing agents requires the use of isolated and purified topoisomerase. Because the assay must be performed using stoichiometric amounts of enzyme, the tester must purify large quantities of enzyme in order to perform a minimal number of tests. Accordingly, there is a need to overproduce the enzyme in a time-conserving and cost-effective manner.

Furthermore, large quantities of enzyme or enzyme fragments may be desired to generate polyclonal or monoclonal antibodies. Such antibodies may then be used in affinity chromatography to facilitate purification of the enzyme. Large quantities of enzyme or enzyme fragments may also be desired to perform X-ray crystallography and other analytical studies.

The present invention solves these problems by identifying, isolating and expressing a type I topoisomerase gene from C. albicans in an expression system suitable for large-scale production. In addition, identification and isolation of a C. albicans type I topoisomerase gene allows for its expression in a variety of systems which facilitate purification of a C. albicans topoisomerase I encoded therefrom. A C. albicans type I topoisomerase gene may also used in a assay system for identifying compounds which inhibit the growth of fungi.

Summary of the Invention

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In one aspect, the present invention provides an isolated and purified single- or double-stranded polynucleotide, typically DNA, having a nucleotide sequence comprising a nucleotide sequence selected from the group consisting of (a) the sense sequence of SEQ ID NO:1 from about nucleotide position 474 to about nucleotide position 2807; (b) a sequence complementary to the sequence of (a); (c) sequences that, when expressed, encode a polypeptide encoded by a sequence of (a); and d) analogous sequences that hybridize under stringent conditions to the sequences of (a) or (b). A polynucleotide of the present invention

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may also comprise the sense sequence of SEQ ID NO:1 from about nucleotide position 345 to about nucleotide position 2807 as well as the longer counterparts of (b), (c) and (d) above. A preferred polynucleotide is a DNA molecule. In another embodiment, the polynucleotide is an RNA molecule.

In another aspect, the present invention provides specific isolated and purified singleor double-stranded fragments of the above mentioned polynucleotide including the fragment counterparts of (b), (c) and (d) above. The specific fragments are from nucleotide position 474 to nucleotide position 923, nucleotide position 2199 to nucleotide position 2624 and nucleotide position 2655 to nucleotide position 2807 as numbered in FIG. 6.

In another embodiment, a DNA molecule of the present invention is contained in an expression vector. The expression vector preferably further comprises an enhancer-promoter operatively linked to the polynucleotide. In an especially preferred embodiment, the DNA molecule in the vector is one of the preferred sequences or fragments mentioned above.

Preferably, a topoisomerase I of the present invention is a recombinant topoisomerase I having the amino acid residue sequence of SEQ ID NO:2 or its subset SEQ ID NO:3. A preferred C. albicans topoisomerase I has about 821 or fewer amino acid residues and comprises the amino acid residue sequence of SEQ ID NO:3. Also useful are the shorter polypeptides encoded by the DNA fragments mentioned above.

In another aspect, the present invention provides a process of making *C. albicans* topoisomerase I comprising transforming a host cell with an expression vector that comprises a polynucleotide of the present invention, maintaining the transformed cell for a period of time sufficient for expression of the topoisomerase I, and recovering the topoisomerase I. Preferably, the host cell is an eukaryotic host cell such as a mammalian, yeast or fungal cell, or a bacterial cell. Especially preferred host cells are *Candida spp.*, *S. cerevisiae* and *E. coli*. The present invention also provides a topoisomerase I made by a process of this invention. A preferred such topoisomerase I is recombinant *C. albicans* topoisomerase I.

In another aspect, the present invention provides for a host cell transformed with a polynucleotide or expression vector of this invention. Preferably, the host cell is a yeast or fungal cell such as S. cerevisiae or Candida spp. or a bacterial cell such as E. coli.

The present invention further provides a method for identifying compounds having the ability to kill or inhibit the growth of fungal strains.

Brief Description of the Drawings

FIG. 1 shows the amino acid sequence used to design sets of degenerate oligonucleotide primers for isolating a 127 base pair (bp) C. albicans DNA fragment with

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homology to eukaryotic topoisomerase I genes. In FIG. 1 and in all Figures hereinafter, the abbreviations "Hu, Sc, Sp, and Dm" have the following meanings: Hu = human, Sc = Saccharomyces cerevisiae, Sp = Schizosaccharomyces pombe, and Dm = Drosophila melanogaster. In FIG. 1, the Hu sequence represents amino acids 589-634 (D'Arpa et al., 1988), the Sc sequence represents amino acids 516-560 (Thrash, et al., 1985), the Sp sequence represents amino acids 561-605 (Uemura, et al., 1987) and the Dm sequence represents amino acids 811-856 (Hsieh, et al., 1992) of each species' respective topoisomerase I.

FIG. 2 shows a comparison of the sequence of the 127 bp C. albicans DNA fragment to nucleotides 2043-2180 of the Saccharomyces cerevisiae topoisomerase I gene.

FIG. 3 shows the predicted alignment of the polypeptide encoded by the 127 bp C. albicans DNA fragment with the amino acid sequences of genes encoding type I topoisomerases from other eukaryotic species. In FIG. 3 and in all Figures hereinafter, the abbreviations "Ca, Um, Mm, XI, and At" have the following meanings: Ca = Candida albicans, Um = Ustilago maydis, Mm = Mus musculus, XI = Xenopus laevis, and At = Arabadopsis thaliana. In FIG. 3, the Ca sequence represents the amino acids encoded by the insert in plasmid pKG1, Sc represents amino acids 514-561 (Thrash et al., 1985), Sp represents amino acids 559-606 (Uemura et al., 1987), Hu represents amino acids 587-635 (D'Arpa et al., 1988), Um represents amino acids 557-616 (Gerhold et al., 1994), Mm represents amino acids 589-637 (Baumgartner et al., 1994), XI represents amino acids 643-691 (Pandit and Sternglanz, 1992), Dm represents amino acids 809-858 (Hsieh et al., 1992), and At represents amino acids 733-780 (Kieber and Signer, 1990) of each species respective topoisomerase I.

FIG. 4 shows the amino acid sequence used to design a set of degenerate oligonucleotide primers for isolating a 1044 bp *C. albicans* DNA fragment with homology to eukaryotic topoisomerase I genes. In FIG. 4, Hu represents amino acids 273-282, *Mm* represents amino acids 275-284, *Dm* represents amino acids 496-505 and *Sc* represents amino acids 201-210.

FIG. 5 shows the sequence of the cloned 1044 bp C. albicans DNA fragment.

FIG. 6(a-j) shows the complete double-stranded sequence of *C. albicans* topoisomerase I gene including upstream and downstream gene sequences and the predicted polypeptide encoded therefrom.

FIG. 7(a-b) shows the comparative homology in amino acids between C. albicans topoisomerase I and human topoisomerase I (d'Arpa, et al., 1988).

FIG. 8 shows the nucleotide sequence of an adapter DNA which carries the 5' and 3' ends of the *C. albicans* topoisomerase I gene.

FIG. 9 shows a schematic diagram of the construction of plasmid pVT100-UCT7-7.

Detailed Description of the Invention

I. The Invention

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The present invention provides isolated and purified polynucleotides that encode a type I topoisomerase from Candida albicans, fragments thereof, expression vectors containing those polynucleotides, host cells transformed with those expression vectors, a process of making C. albicans topoisomerase I using those polynucleotides and vectors, and isolated and purified recombinant C. albicans topoisomerase I and polypeptide fragments thereof. The present invention also provides a method for identifying compounds having antifungal activity using polynucleotides that encode a type I topoisomerase from C. albicans.

II. TOP1 Polynucleotides

In one aspect, the present invention provides an isolated and purified polynucleotide that encodes a type I topoisomerase from the pathogenic fungus, Candida albicans (C. albicans). Hereinafter, the abbreviation "TOP1" is used to designate a gene or polynucleotide sequence encoding a type I topoisomerase; the abbreviation "topo I" is used to designate a polypeptide or an amino acid sequence encoded by a TOP1. A polynucleotide of the present invention that encodes C. albicans topo I is an isolated and purified polynucleotide having a nucleotide sequence which comprises a nucleotide sequence selected from the group consisting of

- (a) the sense sequence of SEQ ID NO:1 from about nucleotide position 345 or 474 to about nucleotide position 2807,
 - (b) a sequence complementary to the sequence of (a),
- (c) sequences that, when expressed, encode a polypeptide encoded by the sequence of (a), and
- (d) analogous sequences that hybridize under stringent conditions to the sequences of (a) or (b). A preferred polynucleotide is a DNA molecule. In another embodiment, the polynucleotide is an RNA molecule.

The nucleotide sequence and deduced amino acid residue sequence of Candida albicans topo I are set forth in SEQ ID NO:1 and SEQ ID NO:2 shown in FIG. 6. The nucleotide sequence of SEQ ID NO:1 is a full-length DNA clone of the gene encoding C. albicans topo I and is intended to represent both the sense strand (shown on top) and its complement (i.e. the strand shown below the top strand in FIG. 6). In eukaryotic cells, translation of RNA initiates at an AUG codon coding for methionine (which, in the C. albicans topo I shown in FIG. 6, corresponds to nucleic acid position 474). However, the

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open reading frame (ORF) of *C. albicans* TOP1 begins at about nucleotide position 345. Thus SEQ ID NO:1 contains two "coding regions" (hereinafter CR1 and CR2) which, in FIG. 6, correspond to nucleotide positions 345-2807 and 474-2807, respectively, either of which may encode the native topo I enzyme. SEQ ID NO:2 is the deduced amino acid residue sequence encoded from CR1 and represents a polypeptide which is 821 amino acids in length. In FIG. 6, SEQ ID NO:2 starts with the amino glutamic acid (E) at position -43 and ends with the amino acid phenylalanine (F) at position 821. SEQ ID NO:3 is the deduced amino acid sequence encoded from CR2 and represents a polypeptide which is 778 amino acids in length. In FIG. 6, SEQ ID NO:3 begins with methionine (M) at position +1 and ends with pheylalanine (F) at position 821.

The present invention also contemplates shorter polynucleotide sequences or fragments of CR1 and CR2 which may or may not have topo I catalytic activity. More specifically, a polynucleotide of the present invention is also an isolated and purified single-or double-stranded polynucleotide having a nucleotide sequence which comprises a nucleotide sequence selected from the group consisting of (a) the sense sequence of SEQ ID NO:1 from about nucleotide position 474 to about nucleotide position to about nucleotide position 923, the sense sequence of SEQ ID NO:1 from about nucleotide position 2199 to about nucleotide position 2624 and the sense sequence of SEQ ID NO:1 from about nucleotide position 2655 to about nucleotide position 2807. The invention is also intended to encompass the fragment counterparts of (b), (c) and (d) mentioned above.

The present invention also contemplates analogous DNA sequences which hybridize under stringent hybridization conditions to the DNA sequences set forth above. Stringent hybridization conditions are well-known in the art and define a degree of sequence identity greater than about 80%-90%. The modifier "analogous" refers to those nucleotide sequences that encode polypeptides having only conservative differences and which retain the conventional characteristics and activities of a topoisomerase I; eg. nicking DNA, changing the DNA's linking number and resealing the nick. The present invention also contemplates naturally-occurring allelic variations and mutations of the DNA sequences set forth above so long as those variations and mutations code, on expression, for a topo I of this invention as set forth hereinafter.

As is well-known in the art, because of the degeneracy of the genetic code, there are numerous other DNA and RNA molecules that can code for the same polypeptides as those encoded by CR1, CR2 and fragments thereof. The present invention, therefore, contemplates those other DNA and RNA molecules which, on expression, encode for the polypeptide of SEQ ID NO:2, its subset SEQ ID NO:3 or fragments thereof. Having identified the amino acid residue sequence encoded by *C. albicans* TOP1, and with knowledge of all triplet codons for each particular amino acid residue, it is possible to

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describe all such encoding RNA and DNA sequences. DNA and RNA molecules other than those specifically disclosed herein, which molecules are characterized simply by a change in a codon for a particular amino acid, are within the scope of this invention.

As is well-known in the art, codons constitute triplet sequences of nucleotides in mRNA molecules and as such, are characterized by the base uracil (U) in place of base thymidine (T) (which is present in DNA molecules). A simple change in a codon for the same amino acid residue within a polynucleotide will not change the structure of the encoded polypeptide. By way of example, it can be seen from SEQ ID NO:1 (see FIG. 6) that a TCA codon for serine (i.e. UCA in mRNA) exists at nucleotide positions 318-320 and again at positions 408-410 and 411-413. However, it can also be seen from that same sequence that serine can be encoded by an AGT codon (see e.g. nucleotide positions 438-440) and a TCG codon (see e.g., nucleotide positions 462-464). Substitution of the latter codons for serine with the TCA codon for serine, or visa-versa, does not substantially alter the DNA sequence of SEQ ID NO:1 and results in expression of the same polypeptide. In a similar manner, substitutions of the recited codons with other equivalent codons can be made in a like manner without departing from the scope of the present invention.

Furthermore, it should be noted that, in *C. albicans*, the CUG codon has been shown to encode serine rather than leucine (see Santos and Tiute, 1995), although this phenomenon is not seen in all *Candida* species or in other fungi (such as *S. cerevisiae*). Thus, a recombinant *C. albicans* TOP1 of the present invention expressed in a *C. albicans* strain would most likely encode a topo I having a serine residue at amino acid position 407 (Xaa) in SEQ ID NO:2 and 321 in SEQ ID NO:3 (corresponding to the CUG at nucleotide position 1434 and amino acid "X" at position 321 in SEQ ID NO:1). In contrast, expression of *C. albicans* TOP1 in another *Candida* or other fungal species could result in placement of a leucine residue at this same position. Thus, the present invention contemplates a topo I having either a serine or leucine residue at the Xaa position depending on the expression system used.

A polynucleotide of the present invention can also be an RNA molecule. A RNA molecule contemplated by the present invention is complementary to or hybridizes under stringent conditions to any of the DNA sequences set forth above. Exemplary and preferred RNA molecules are mRNA molecules that encode a topo I of this invention.

The present invention also contemplates oligonucleotides from about 15 to about 50 nucleotides in length, which oligonucleotides serve as primers and hybridization probes for the screening of DNA libraries and the identification of DNA or RNA molecules that encode *C. albicans* topo I. Such primers and probes are characterized in that they will hybridize to polynucleotide sequences encoding topo I. An oligonucleotide probe or primer contains a nucleotide sequence of at least 15 nucleotides that is identical to or complementary to a

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contiguous sequence of a topoisomerase I polynucleotide of the present invention. Thus, where an oligonucleotide probe is 25 nucleotides in length, at least 15 of those nucleotides are identical or complementary to a sequence of contiguous nucleotides of a TOP1 of the present invention. Exemplary TOP1 polynucleotides of the present invention are set forth above.

An oligonucleotide primer or probe of the present invention can be prepared using standard procedures well-known in the art. A preferred method of polynucleotide synthesis is via cyanoethyl phosphoramidite chemistry. A detailed description of the preparation, isolation and purification of polynucleotides encoding *C albicans* topo I is set forth in the Examples below.

III. C. albicans topo I Polypeptides

In another aspect, the present invention provides a recombinant type I topoisomerase from C albicans. A C. albicans topo I of the present invention is a polypeptide of about 821 or fewer amino acid residues. As set forth in FIG. 7, forms of type I topoisomerases have been identified in various species with from 314 to 972 amino acid residues. The various forms of topo I are characterized by a moderate degree of sequence identity. By way of example, the identity between human and C. albicans enzyme is 45% at the amino acid level.

When the amino acid residue sequence encoded by *C. albicans* TOP1 was compared against other known amino acid residue sequences using a database searching algorithm, only moderate sequence similarity was found. The varying degree of identity among type I topoisomerases from various species allows for definition of the *C. albicans* topo I amino acid residue sequence by the regions lacking identity. Thus, in one embodiment, a *C. albicans* topo I is an isolated and purified polypeptide of about 821 or less amino acid residues, comprising at least one of the following amino acid residue sequences:

- a) from residue position 1 to residue position 150 of SEQ ID NO:3;
- b) from residue position 297 to residue position 529 of SEQ ID NO:3;
- c) from residue position 573 to residue position 717 of SEQ ID NO:3; and
- d) from residue position 728 to residue position 778 of SEQ ID NO:3.

Preferably, a topo I of the present invention comprises two or more of the above sequences. Most preferably, the topo I has all of the above sequences.

More preferably, a topo I of the present invention is a recombinant topo I from C. albicans. Although generally, eukaryotic polypeptides are known to begin with a methionine residue, the present invention also contemplates a recombinant topo I encoded by the complete ORF of C. albicans TOP1. Thus, recombinant C. albicans topo I can be defined as a polypeptide of about 821 or less amino acid residues comprising the amino acid

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residue sequence of SEQ ID NO:2. A preferred recombinant C. albicans topo I has the amino acid residue sequence of SEQ ID NO:3.

The present invention also contemplates amino acid residue sequences that are substantially duplicative of the sequences set forth herein such that those sequences demonstrate like biological activity when compared to disclosed sequences. Such contemplated sequences include those sequences characterized by a minimal change in amino acid residue sequence or type (e.g., conservatively-substituted sequences) which insubstantial change does not alter the fundamental nature and biological activity of *C. albicans* topo I.

It is well known in the art that modifications and changes can be made in the structure of a polypeptide without substantially altering the biological function of that peptide. For example, certain amino acids can be substituted for other amino acids in a given polypeptide without any appreciable loss of function. In making such changes, substitutions of like amino acid residues can be made on the basis of relative similarity of side-chain substituents, for example, their size, charge, hydrophobicity, hydrophilicity, and the like.

As detailed in United States Patent No. 4,554,101, incorporated herein by reference, the following hydrophilicity values have been assigned to amino acid residues: Arg (+3.0); Lys (+3.0); Asp (+3.0); Glu (+3.0); Ser (+0.3); Asn (+0.2); Gln (+0.2); Gly (0); Pro (-0.5); Thr (-0.4); Ala (-0.5); His (-0.5); Cys (-1.0); Met (-1.3); Val (-1.5); Leu (-1.8); Ile (-1.8); Tyr (-2.3); Phe (-2.5); and Trp (-3.4). It is expected that an amino acid residue can be substituted for another having a similar hydrophilicity value (e.g., within a value of plus or minus 2.0) and still result in a biologically equivalent polypeptide.

In a similar manner, substitutions can be made on the basis of similarity in hydropathic index. Each amino acid residue has been assigned a hydropathic index on the basis of its hydrophobicity and charge characteristics. Those hydropathic index values are: Ile (+4.5); Val (+4.2); Leu (+3.8); Phe (+2.8); Cys (+2.5); Met (+1.9); Ala (+1.8); Gly (-0.4); Thr (-0.7); Ser (-0.8); Trp (-0.9); Tyr (-1.3); Pro (-1.6); His (-3.2); Glu (-3.5); Gln (-3.5); Asp (-3.5); Asn (-3.5); Lys (-3.9); and Arg (-4.5). In making a substitution based on the hydropathic index, a value of within plus or minus 2.0 is preferred.

A topo I of the present invention has numerous uses. By way of example, such a polypeptide can be used in assays which identify compounds that stabilize the cleavage complex of topo I and DNA. A topo I polypeptide can also be used to design compounds that interact with the cleavage complex of topo I and DNA. More specifically, the topo I or a portion of the topo I can be crystallized and used for designing drugs that stabilize the cleavage complex formed between the topo I and DNA.

In addition, a topo I of the present invention or fragments thereof may used to

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produce antibodies that immunoreact specifically with *C. albicans* topo I. Such antibodies may then be used to isolate and purify *C. albicans* topo I. Means for producing antibodies are well known in the art. An antibody directed against *C. albicans* topo I can be a polyclonal or a monoclonal antibody.

Antibodies against *C. albicans* topo I can be prepared by immunizing an animal with a topo I polypeptide of the present invention. Means for immunizing animals for the production of antibodies are well-known in the art. By way of an example, a mammal can be injected with an inoculum that includes a polypeptide as described herein above. The polypeptide can be included in an inoculum alone or conjugated to a carrier protein such as BSA or keyhole limpet hemocyanin (KLH). The polypeptide can be suspended, as is well-known in the art, in an adjuvant to enhance the immunogenicity of the polypeptide. Sera containing immunologically active antibodies are then produced from the blood or other fluid (eg. ascites) of such immunized animals using standard procedures well-known in the art.

The identification of antibodies that immunoreact specifically with *C. albicans* topo I is made by exposing sera suspected of containing such antibodies to a polypeptide of the present invention to form a conjugate between antibodies and the polypeptide. The existence of the conjugate is then determined using standard procedures well-known in the art.

A topo I polypeptide of the present invention or fragments thereof may also be used to prepare monoclonal antibodies against topo I and used as a screening assay to identify such monoclonal antibodies. Monoclonal antibodies are produced from hybridomas prepared in accordance with standard techniques such as that described by Kohler *et al.*. Briefly, a suitable mammal (e.g., BALB/c mouse) is immunized by injection with a polypeptide of the present invention. After a predetermined period of time, splenocytes are removed from the mouse and suspended in a cell culture medium. The splenocytes are then fused with an immortal cell line to form a hybridoma. The formed hyridomas are grown in cell culture and screened for their ability to produce a monoclonal antibody against topo I. Screening of the cell culture medium is made with a polypeptide of the present invention.

IV. Method of Making C. albicans topo I.

In another aspect, the present invention provides a process of making *C. albicans* topo I. In accordance with that process, a suitable host cell is transformed with a polynucleotide of the present invention. The transformed cell is maintained for a period of time sufficient for expression of the topo I; the topo I is then recovered.

Means for transforming host cells in a manner such that those cells produce recombinant polypeptides are well-known in the art. Briefly, a polynucleotide that encodes

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the desired polypeptide is placed into an expression vector suitable for a given host cell. That vector can be a viral vector, phage or plasmid and may be one which either replicates autonomously within the host cell or integrates into the host cell chromosome. In a preferred embodiment, a host cell used to produce topo I is an eukaryotic host cell and an expression vector is an eukaryotic expression vector (i.e., a vector capable of directing expression in a eukaryotic cell). Such eukaryotic expression vectors are well known in the art. An especially preferred eukaryotic host cell is an S. cerevisiae or C. albicans. A preferred expression vector is a vector capable of directing expression in S. cerevisiae or C. albicans.

In another preferred embodiment, the host cell is a bacterial cell. An especially preferred bacterial cell is an $E.\ coli.$ Thus, a preferred expression vector is a vector capable of directing expression in $E.\ coli.$

A polynucleotide of an expression vector of the present invention is preferably operatively associated or linked with an enhancer-promoter. A promoter is a region of a DNA molecule typically within about 100 nucleotide pairs in front of (upstream of) the point at which transcription begins. That region typically contains several types of DNA sequence elements that are located in similar relative positions in different genes. As used herein, the term "promoter" includes what is referred to in the art as an upstream promoter region or a promoter of a generalized RNA polymerase transcription unit. Preferred promoters are those which are operative in bacteria such as a *lac* promoter. Even more preferred promoters are those which are operative in yeast or fungi. Especially preferred promoters are an ADH1 or GAL 10 promoter of *S. cerevisiae* or a *C. albicans* promoter.

Another type of transcription regulatory sequence element is an enhancer. An enhancer provides specificity of time, location and expression level for a particular encoding region (e.g., gene). A major function of an enhancer is to increase the level of transcription of a coding sequence in a cell that contains one or more transcription factors that bind to that enhancer. Unlike a promoter, an enhancer can function when located at variable distances from a transcription start site so long as the promoter is present.

As used herein, the phrase "enhancer-promoter" means a composite unit that contains both enhancer and promoter elements. An enhancer-promoter is operatively linked to a coding sequence that encodes at least one gene product. As used herein, the phrase "operatively linked" or its grammatical equivalent means that a regulatory sequence element (e.g. an enhancer-promoter or transcription-terminating region) is connected to a coding sequence in such a way that the transcription of that coding sequence is controlled and regulated by that enhancer-promoter. Means for operatively linking an enhancer-promoter to a coding sequence are well-known in the art.

An enhancer-promoter used in an expression vector of the present invention can be any enhancer-promoter that drives expression in a host cell. By employing an enhancer-

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promoter with well-known properties, the level of expression can be optimized. For example, selection of an enhancer-promoter that is active in specifically transformed cells permits tissue or cell specific expression of the desired product. Still further, selection of an enhancer-promoter that is regulated in response to a specific physiological signal can permit inducible expression.

A coding sequence of an expression vector is operatively linked to a transcription-terminating region. RNA polymerase transcribes an encoding DNA sequence through a site where polyadenylation occurs. Typically, DNA sequences located a few hundred base pairs downstream of the polyadenylation site serve to terminate transcription. Those DNA sequences are referred to herein as transcription-termination regions. Those regions are required for efficient polyadenylation of transcribed messenger RNA (mRNA). Enhancer-promoters and transcription-terminating regions are well-known in the art. The selection of a particular enhancer-promoter or transcription-terminating region will depend, as is also well-known in the art, on the cell to be transformed.

The present invention also comtemplates the use of expression vectors which facilitate purification of a desired polypeptide. For example, a gene encoding the desired polypeptide may be cloned into an expression vector which, when expressed, produces the polypeptide fused or linked to a chemical or biological tag. A tag may be any chemical or biological compound or fragment thereof capable of binding to a specific substrate or receptor. Thus, tags serve to facilitate purfication of a fusion product (i.e. tag/polypeptide) via specific binding of the tag portion to its receptor or substrate. Preferably the tag is fused to the polypeptide in a manner that permits it to be cleaved from the polypeptide after purification. In illustration, C. albicans TOP1 could be cloned into a pGEX vector (Pharmacia Biotech, Inc., Piscataway, New Jersey) which, when placed in a suitable host, would express a fusion protein of C. albicans topo I and glutathione S-transferase (GST). The fusion protein could then be purified by affinity chromatography using glutathione sepharose 4B (which binds to the GST portion of the fusion product) and the topo I enzyme cleaved from the GST tag using a site-specific protease, the recognition sequence of which is located upstream from the topo I. Other commercially available tags could also be used and are well-known to those of ordinary skill in the art.

The present invention also contemplates a host cell transformed with a polynucleotide or expression vector of this invention. Means for transforming cells and polynucleotides and expression vectors used to transform host cells are set forth above. Preferably, the host cell is an eukaryotic host cell such as a yeast or fungal cell or a prokaryotic cell such as an *E. coli*.

V. Method of Identifying Stabilizing Agents of C. albicans topo I and DNA Cleavage

Complex

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In another aspect, the present invention provides a method for identifying antifungal compounds and, more specifically, antifungal compounds which act as stabilizing agents of a C. albicans topo I and DNA cleavage complex. A "cleavage complex", as used herein, refers to a transient association between a topoisomerase and DNA during which time the enzyme performs various catalytic functions (such as binding to and nicking a particular region of the DNA, passing a DNA segment through or around the break to change the linking number, and ultimately resealing (i.e. repairing) the nick). The term "stabilizing agent" as used herein refers to a chemical compound which inhibits cell growth or promotes cell death through some manner of interaction with a topoisomerase I/DNA cleavage complex. The term "stabilize," or "stabilization" when used herein and in reference to a stabilizing agent and a cleavage complex, refers to that interaction of the stabilizing agent with the cleavage complex which leads to significant inhibition of cell growth and reproduction or, preferably, cell death. As shown by Nitiss and Wang (1988), the chemical compound camptothecin stabilizes the topo I/DNA cleavage complex of Saccharomyces cerevisiae, inducing fungal death. Notwithstanding this observation, the nature of the interaction between the stabilizing agent and the cleavage complex and the mechanism by which the agent induces cell death is unknown.

The present invention provides an assay method for identifying stabilizing agents of a C. albicans topo I/DNA cleavage complex. Assays for identifying stabilizing agents may be carried out both in whole-cell preparations and in ex vivo cell-free systems. In each instance, the assay target is a C. albicans topoisomerase I/DNA cleavage complex, wherein the C. albicans topoisomerase I is a C. albicans topo I of the present invention. It is expected that the assay methods of the present invention will be suitable for both small- and large-scale screening of test compounds, as well as in quantitative assays such as serial dilution studies wherein the target topo I/DNA cleavage complex is exposed to a range of test compound concentrations.

When the method of the present invention is carried out as a whole-cell assay, the target is an intracellular topo I/DNA cleavage complex and the entire, living fungal cell is exposed to the test compound under conditions normally suitable for growth. Such conditions, including essential nutrients, optimal temperatures and other parameters, depend upon the particular fungal strain being used and are well-known in the art. Stabilization of the topo I/DNA cleavage complex may be determined by observing the cell culture's growth or lack thereof; such observation may be made visually, by optical densitometric or other light absorption/scattering means, or by yet other suitable means, whether manual or automated.

In the above whole-cell assay, an observed lack of cell growth may be due to

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stabilization of the target topo I/DNA cleavage complex or due to an entirely different effect of the test compound so that further evaluation is required to establish the mechanism of action and to determine whether the test compound is a specific stabilizing agent of the cleavage complex. Accordingly, and in a preferred embodiment of the present invention, the method may be performed as a paired-cell assay in a manner similar to that described by Housey in U.S. Patent No. 4,980,281, published December 25, 1990. In a paired-cell assay of the present invention, each test compound is separately tested against two different fungal strains, one having greater topo I activity than the other. Most preferably, such strains are isogenic to each other with respect to this particular trait, meaning that they are equivalent genetically except in their expression of TOP1. (By way of example, isogenic strains may differ with respect to the TOP1 gene by having a copy of a TOP1 gene not found in the other strain or by having multiple copies of a TOP1 gene wherein the other strain has only one copy). In this circumstance, those cells of a strain having greater topo I activity will be more susceptible to the lethal effects of a stabilizing agent than an isogenic strain of cells having lower topo I activity and thus will have comparatively greater mortality or inhibition of cell growth.

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For preparing the above described paired-cell assay, one strain of cells may be altered by any manner of chemical or genetic modification to produce a modified strain having differential topoisomerase I activity relative to a second strain. For example, cells may be chemically modified by exposure to mutagenic agents (such as carcinogenic compounds or UV radiation) to produce a modified strain which underexpresses TOP1 relative to the unmodified parent. In this case, it is possible that such a strain and its parent would not be isogenic to each other since random chemical mutagenesis could affect other genes in addition to a TOP1 gene. However, a chemically modified strain could be further modified by genetic manipulation (to remove extraneous genetic changes by performing crosses to the parent strain and recovering progeny having reduced topo I) for use in a paired-cell assay; in such a case, the parent (i.e. without the additional chemical modification) would serve as an isogenic control. Alternatively, a TOP1 gene may be chemically modified ex vivo by exposure to a mutagenic agent and then reintroduced to a cell to produce a modified cell which expresses altered topo I activity relative to its unmodified parent. In yet another example, the activity of a topo I can be modified in vivo by the presence of a DNA binding agent (such as ethidium bromide) which can be introduced into the culture medium of one strain to alter the topo I activity of that strain relative to the same strain grown without the binding agent. Thus, the initial condition of the cells is not critical to the practice of a paired-cell assay; one need only be able to modify the cells in some manner that is not lethal and which generates a measurable difference in topo I activity.

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A strain may also be genetically manipulated to alter the production of topo I in a particular cell. For example, a strain may be modified to either produce or over-produce topo I by cloning a *C. albicans* TOP1 of the present invention into a plasmid or suitable expression vector and transforming the cloned vector into a suitable host strain by means known to those of ordinary skill in the art. Suitable expression vectors and host cells are those previously described in Section IV above. Other preferred expression vectors are the non-integrative double-ARS shuttle vectors described by Pla *et al.* (1995) which are able to autonomously replicate in *C. albicans*.

For determining whether a compound acts as a fungicide or inhibitor of fungal growth, both the modified strain (harboring an expression vector comprising a TOP1 gene of the present invention) and the isogenic parental strain (harboring the expression vector only) would be exposed simultaneously to a compound of interest under essentially identical growth conditions. A compound which induces death of the modified strain or which inhibits that strain's growth would be a putative stabilizing agent.

Alternatively, a strain may be genetically modified to reduce the activity of a native topo I by any means known to disrupt a TOP1 coding sequence. For example, a cloned C. albicans TOP1 of the present invention can be used to disrupt or inactivate a native TOP1 within a host strain (i.e. in vivo). Techinques for performing gene disruption are well known to those of ordinary skill in the art, (see for example, Fonzi, W.A. and Irwin, M.Y., 1993 and Rothstein, R., 1991). As one example, targeted integration of a selectable marker into the chromosomal copy of a yeast or fungal TOP1 may be accomplished by first cloning the selectable marker gene into a single site within a coding sequence of C. albicans TOP1 (i.e. CR1 or CR2) or a fragment thereof. Selectable marker genes are those which confer upon a cell the ability to grow under a particular set of growth conditions (i.e. in selective medium). Representative examples of selective marker genes include but are not limited to genes which confer drug resistance, such as G418 and genes which confer upon a cell the ability to grow in the absence of an exogenously supplied essential nutrient.

When a fragment of a coding region is used, it should be of a such a length that the marker gene, after cloning, is flanked on each side by a sequence of the coding region that is at least about 250 base pairs in length. The cloned fragment is then transformed into the targeted host cell to induce recombination into the chromosomal copy, and then grown on selective media to identify recombinant cells. As a consequence, functional topo I will not be produced by that particular cell.

As another example, a non-functional topo I may be generated by genetically altering a chromosomal copy of a TOP1 through the process of gene replacement. Gene replacement may be accomplished by specifically altering the DNA sequence of a C. albicans TOP1 (so as to render the topo I encoded therefrom non-functional), subcloning the

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altered sequence into a suitable vector capable of recombining in the yeast or fungal host, and introducing the vector containing the subcloned sequence into the appropriate host so that exchange of the wild-type allele with the mutated one will occur. The steps involved in gene replacement employ standard recombinant DNA techniques and are well known to those of ordinary skill in the art.

To identify stabilizing agents using modified cells having reduced or no topo I activity, both the modified strain and the parental strain would be exposed to a compound or compounds of interest under conditions suitable for growth. In this case however, the parental strain would be most susceptible to a stabilizing agent and would die or show inhibited growth as a result of exposure to the compound.

Any techniques for directly or indirectly measuring topo I protein or topo I activity are suitable for establishing differential topo I production between a modified strain and its parent. For example, quantitative analysis of topo I may be performed by preparing lysates of cells, partially purifying the topo I from the lysates and measuring topo I activity ex vivo (cf. Fostel et al., 1992). Alternatively, when a modified strain is generated by the introduction of a multicopy plasmid containing a TOP1, the actual copy number of the plasmid in the modified strain may be quantified. As another example, a modified strain and its parent may be preliminarily tested for sensitivity to a known stabilizing agent such as camptothecin. Thus, the technique selected may vary depending upon the manner in which the modification is achieved but such techniques are either well known to those of ordinary skill in the art or are taught in the present specification.

The invention will be better understood in connection with the following examples, which are intended as an illustration of and not a limitation upon the scope of the invention. Both below and throughout the specification, it is intended that citations to the literature are expressly incorporated by reference.

EXAMPLE 1 Identification of C. albicans topo I Gene

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1a. <u>Isolation of a 127 bp DNA fragment from C. albicans</u> having homology to eukaryotic type 1 topoisomerase genes

Genomic DNA was isolated from C. albicans strain ATCC 10321, by the method of Polaina and Adam (1991). Cells were grown in YEPD broth (1% Bacto yeast extract, 2% Bacto peptone, 2% glucose, all from Difco, Detroit, MI) to a density of approximately 8 absorbance units, harvested, washed in TGE (20 mM KCl, 1 mM EDTA, 20 mM Tris pH

7.4, 10% glycerol), and frozen at -70°. Cells were thawed, resuspended in lysis buffer (3% sodium lauryl sarcosinate (SDS), 10 mM Tris-HCl pH 7.5, 1 mM EDTA), extracted with phenol/chloroform and precipitated with ethanol. The resulting DNA was used as a template for the polymerase chain reaction (PCR; Saiki et al., 1988). Two sets of degenerate primers for PCR were selected based on conserved regions in other eukaryotic type I topoisomerases (Hsieh et al., 1992) as shown in FIG. 1. New sites for restriction endonucleases Xba I and Bam H1 were also introduced into the primers to facilitate subcloning. The primer sets used were:

5'-CTCTAGATTYMGNACNTAYAAYGC-3' SEQ ID NO:4:

5'- TGGATCCYTGRTGRTTRCANARDAT-3' SEQ ID NO:5:

The symbols shown in these primers (and in all other primers mentioned herein) have the meanings shown in Table 2.

TABLE 2

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Symbol	Base			
Α	Adenine			
C	Cytosine			
D	any base but cytosine (not C)			
G	Guanine			
M	A or C only			
N	any base			
R	A or G only			
T	Thymine			
U	Uracil			
Y	C or T/U only			

The PCR reaction was carried out using reagents from Perkin Elmer (Norwalk, CN), according to the manufacturer's recommendations, using 2 µg genomic C. albicans DNA and approximately 1 nmole of each of SEQ ID NO:4 and SEQ ID NO:5 (synthesized by Northwestern University, Evanston, IL). The reactions were incubated at 45°C for 1 minute, raised to 72°C over a two minute interval, held at 72°C for 1 minute, then at 94°C for 1 minute. This pattern was repeated for 50 cycles. The resulting DNA products were separated using a TBE-urea 17-26% polyacrylamide gel (Enprotech, Natick, MA); each band was characterized according to the following method:

The DNA band of interest was excised from the gel using the "crush and soak method" (Sambrook et al., 1982). Briefly, the fragment of gel containing the DNA of interest was placed in a tube, crushed with a sterile pipette, and incubated at 37°C for 16 hours in elution buffer (0.5 M ammonium acetate, 10 mM magnesium acetate, 0.1% SDS, 1 mM EDTA, pH 8.0). The excised DNA was incubated with the Klenow fragment of DNA

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polymerase I (Boehringer Mannheim, Indianapolis, IN) in the presence of the four nucleotide triphosphates to fill in single-stranded termini, then extracted with phenol/chloroform and precipitated with ethanol. Plasmid pBSK+ (Stratagene GmbH, Heidelberg, Germany) was cut with EcoRV (GIBCO BRL, Gaithersburg, MD). This and other restriction digests were performed in the manufacturer's recommended reaction buffer, and unless otherwise indicated, all restriction and modifying enzymes were obtained from this same source. After digestion, the plasmid DNA was extracted with phenol/chloroform and precipitated with ethanol. The excised DNA was ligated to the plasmid using T4 DNA ligase in the manufacturer's reaction buffer, at 17° C for 16 hours. The resulting DNA was used to transform competent Maximum Efficiency DH5\alphaF'IQ E. coli (GIBCO BRL, Gaithersburg, MD), following the manufacturer's recommended protocol, and an aliquot of 100 µL of the cells was plated on LB-agarose (Lennox broth; from Micro Diagnostics, Lombard, IL) containing 150 µg/mL ampicillin. Prior to use, 35 µL of a 50 mg/mL X-Gal (5-bromo-4-chloro-3-indolyl- β -D-galactoside; Sigma, St. Louis MO) solution in dimethyl formamide and 20 μL of a 100 mM IPTG (Isopropyl β-D-thiogalactopyranoside; Sigma, St. Louis, MO) solution in water were spread on the plate. DNAs from clones carrying inserts were identified on the basis of altered mobility on agarose gels.

The sequence of the inserts was determined using commercial primers from Stratagene. The primers,

5'-AATTAACCCTCACTAAAGGG-3'

SEQ ID NO:6

5'-GTAATACGACTCACTATAGGGC-3'

SEQ ID NO:7

are homologous to the T3 and T7 promoter sequences respectively, in the pBSK+ plasmid. Primers were end-labeled with $\gamma^{32}P$ using polynucleotide kinase (Boehringer Mannheim, Indianapolis, IN) in the buffer provided by the manufacturer, then extended with Taq DNA polymerase (Boehringer Mannheim, Indianapolis, IN) using dideoxy-sequencing reagents and the method obtained from Promega (Madison, WI). The sequence of the insert from one clone designated 6AB was determined and is given in FIG. 2 (see "Ca"). This insert sequence was also compared to a yeast TOP1, i.e. to nucleotides 2034-2180 of the Saccharomyces cerevisiae TOP1 (see sequence "Sc" in FIG. 2) using the BESTFIT program (GCG, Wisconsin Software package, Madison WI) and the DNA Strider 1.2 program (C. Marck, Centre d'Etudes de Saclay, Gif-Sur-Yvette Cedex, France).

The pBSK+ plasmid with the 6AB fragment inserted was renamed pKG1. A BLAST search (GCG, Wisconsin Software package, Madison WI) of DNA sequences in the GENEMBL database identified other topoisomerase genes as those sequences with the highest score (i.e. having the greatest percent homology to the 6AB insert). An alignment of the predicted *C. albicans* protein sequence encoded by 6AB with other homologous regions of type I topoisomerases from other species is shown in FIG. 3.

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Hybridization to *C. albicans* genomic DNA using the method of Southern (1975) revealed the presence of a single copy sequence homologous to pKG1.

1b. <u>Isolation of a 1044 bp DNA fragment having homology to eukaryotic type I</u> topoisomerase genes

A new set of degenerate primers, based on the sequence

5'-AARAAYTTYTTYAARGA-3' (SEQ ID NO:8) was constructed for the purpose of synthesizing an additional portion of the *C. albicans* topo I gene via PCR. The design of SEQ ID NO:8 was based on anticipated homology to other eukaryotic type I topoisomerases as shown in FIG. 4. SEQ ID NO:8 and a homologous primer,

5'-AGCCACCGTTCTATTGGCAGC-3' (SEQ ID NO:9) derived from the 6AB sequence, were used in a PCR reaction with genomic *C. albicans* DNA as template. The reaction was carried out using reagents from Perkin Elmer, according to the manufacturer's recommendations, with 2 μg genomic *C. albicans* DNA and approximately 1 nmole of each primer. The reactions were incubated at 45° C for 2 minutes, raised to 72° C over a one minute interval, held at 72° C for 3 minutes, then 94° for 1 minute. This pattern was repeated for 50 cycles. A DNA species having the anticipated size of approximately 1 kb was produced.

The PCR products were electrophoresed on an 0.8% low melt agarose gel (LMPA, BRL, Gaithersberg, MD), and the gel slice containing the DNA of interest was excised, melted and subcloned directly into pGEM-T TM (Promega, Madison, WI) using the manufacturer's recommended conditions. The resulting plasmid was verified by restriction analysis, and termed pAT100.

The insert of C. albicans DNA in pAT100 was sequenced using a fmol DNA sequencing kit obtained from Promega (Madison, WI), with primers homologous to the T3 and M13 sequences in the vector, i.e. SEQ ID NO:6 and SEQ ID NO:10 (5'-GTAAAACGACGCCAGT-3'). The insert sequence obtained (SEQ ID NO:11) is given in FIG. 5.

1c. Comparative homology of the 1044 bp DNA fragment to other eukaryotic type 1 topoisomerase genes

The 1044 bp sequence was used as a probe for a search of the GENEMBL (European Molecular Biology Organization) sequence database using the BLAST routine (GCG Wisconsin software package, University of Wisconsin, Madison, WI). BLAST uses the algorithm of Altschul et al. (J. Mol. Biol. 215: 403-410 (1990)) to search the database entries

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for similarity to the probe sequence. The results showed that the highest scoring hits of the insert sequence were to other eukaryotic type I topoisomerases, suggesting that the 1044 bp fragment was indeed part of a *C. albicans* topoisomerase I gene.

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EXAMPLE 2 <u>Isolation of C. albicans topo I gene</u>

2a. Construction of a C. albicans Genomic Library

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C. albicans genomic DNA was prepared according to Olson et al. (1979). Cells were grown overnight in 500 mL of Mock YEPD liquid medium (purchased from BIO101, Vista, CA) from a 3 mL inoculum. The culture was divided into four equal volumes, harvested by centrifugation and washed with water two times. Approximately 6 g wet weight of cells were harvested in total. The 1.5 g aliquots of cells were then resuspended in 4 mL of SCE (1.0 M sorbitol, 0.1 M sodium citrate, 0.06 M EDTA, pH 7) to which was added 2 mg of zymolyase 5000 and 30 µL of \(\beta\)-mercaptoethanol (Sigma, St. Louis, MO) to cause the formation of spheroplasts by degradation of the cell wall. Spheroplasting was followed microscopically by treating an aliquot of cells with 10% SDS. Spheroplasted cells were then added to 7 mL of lysis solution (3% SDS, 0.5 M Tris-HCL. pH 9.0, 0.2 M EDTA), incubated at 65° C for 15 minutes, and cooled rapidly to room temperature. The lysates were layered on top of 50/20/15% sucrose solutions which also contained 0.8 M NaCl, 20 mM Tris-HCL pH 8.0, and 10 mM EDTA. Gradients were centrifuged at 26,000 rpm for 2.5 hours at 20°C using a Beckman SW28 rotor. DNA was collected from the top of the 50% sucrose pad, extracted by rolling very gently with phenol/chloroform for 20 minutes, dialyzed against TE buffer (10mM Tris HCl pH 7.5, 5.0 mM EDTA) and then concentrated against solid sucrose.

Approximately 200 μ g of this *C. albicans* genomic DNA was used by Stratagene (La Jolla, CA) to construct a genomic DNA library in λ FIX[®]II (Stratagene, La Jolla, CA). The λ FIX[®]II vector accepts inserts of size 9-20 kb, so assuming an average insert size of 15 kb, approximately 1000 clones would be expected to contain one genome equivalent of *C. albicans* DNA (genome size 15 megabases). This range of insert size was also considered to be the most useful size for isolation of a full length gene product from *C. albicans*, as the topoisomerase I gene would be expected to span at least 2.5 kb, assuming similarity to the *S. cerevisiae* TOP1 gene. The *C. albicans* genomic DNA was partially digested with Sau3A and size-selected by gel electrophoresis. It was then ligated into the λ FIX[®]II vector following digestion with Xho I and treatment to a partial fill-in reaction, then grown in a P2

host (XL-1 Blue MRA(P2)). Both unamplified and 1x amplified libraries prepared according to this procedure were obtained from Stratagene.

2b. <u>Identification of Phage with Homologous DNA Insert</u>

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An aliquot of the lx amplified *C. albicans* genomic library was used to infect XL1-Blue MRA (P2) according to the manufacturer's suggested conditions. Approximately 25,000 plaques were plated on NZCYM agar (GIBCO, BRL, Gaithersburg, MD) and incubated for 8 hours at 37 °C. Three plates were prepared, and the protocol given by Stratagene was followed to process the plaques for subsequent screening. Filters containing the plaques were prehybridized in PIPES buffer, 50% formamide, 100 µg salmon sperm DNA and 0.5% SDS. The filters were prehybridized at 42 °C for 2 hours followed by hybridization at 42 °C for 24 hours with a probe prepared as described below.

The probe was prepared for hybridization by first digesting pAT100 with Pst1 and Sph1. The digested DNA was electrophoresed on an 0.8% low melt agarose gel (LMPA, GIBCO BRL, Gaithersburg, MD), the 1 kb band was excised and labeled with γ^{32} P-dNTP using a Mega Prime labeling kit obtained from Amersham (Arlington Heights, IL). After hybridization, the filters were washed two times with 2X SSC for 15 minutes at 65°C (1X SSC is 0.15M NaCl, 0.015M Na citrate) followed by washes for 30 minutes at 65°C with 2X SSC, 0.1% SDS and then 0.2X SSC, 0.1% SDS. Filters were air dried and exposed at -70°C to Kodak X-AR tm film (Eastman Kodak, New Haven, CT). Films were developed according to manufacturer's recommended conditions.

Thirty positive plaques from a total of approximately 45 were purified by one round of plaque purification using the labeled probe derived from pAT100. Two of the plaques yielding the strongest hybridization signal, #3 and #26, were arbitrarily selected for a third round of plaque purification following which DNA was prepared from clone #26 using a Wizard Lambda DNA Purification kit from Promega (Madison, WI).

EXAMPLE 3

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Characterization of C. albicans topo I gene

3a. Sequencing Methodology

New sequencing primers were prepared to the 5' and 3' ends of the pAT100 insert in order to read out from each end of the sequence into the remainder of the gene. Sequencing by the dideoxy method of Sanger et al. (1977) was performed on the lambda templates directly, with the use of a fmol DNA sequencing kit from Promega. As new sequence was

determined, new primers were synthesized and used to 'walk' along the clones to obtain a complete open reading frame (ORF). Sequencing on the opposite strand using another set of primers confirmed the sequence. A complete sequence was considered to be obtained following translation of the ORFs and comparison to *S. cerevisiae* and *Schizosaccharomyces pombe* topo I gene sequences. TRANSLATE, BESTFIT, PILEUP and BLAST analyses (GCG, Wisconsin Software package, UW Madison, WI) were used in this determination. The primers used for the sequence determination are given in Table 3, and are highlighted in bold type in FIG. 6.

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TABLE 3

	Primer No.	Nucleotide Position FIG. 6	on Sequence	SEQ ID NO:
15	1	56-76	5'-GCTCCGCACATTCTATTACACC-3'	12
	2	353-376	5'-CCATCAATTTACTCATCAGTTTGG-3'	13
	3	624-646	5'-GAAGAAGATGAAGACGAAGTCCC-3'	14
	4	898-918	5'-GGTGGGAAGTGAATCAAGAAG-3'	15
	5	857-879	5'-GGTTAAATCTGAAACTCCTGAAG-3'	16
20	6	1134-1155	5'-GATTTTTTGGAAGTTTTAAAAG-3'	17
	7	1200-1221	5'-GATTTTAGTAAAATGTATGCTC-3'	18
	8	1410-1428	5'-GGGAAATTGAAACGTCGAG-3'	19
	9	2074-2096	5'-CCATGCAAGATCAAATTGATATA-3'	20
	10	2286-2307	5'-GAACCCAAATTGAAAAAGAAAG-3'	21
25	11	2442-2466	5'-GGTAAACCATTATTAACTGAATCAG-3	• 22
	12	2617-2640	5'-CAATTCAATTAAAAGATAAAGAAG-3'	23
	13	2870-2891	5'-GTACCTGTAATATATCCAC-3'	24*
	14	2914-2937	5'-CCTTTCACACTGCAGTATCAATC-3'	25*
	15	2800-2819	5'-CCCCTAATTTTAGAATCTCC-3'	26
30	16	2640-2662	5'-CCTAAAGAAACTTCAGAATTATC-3'	27
	17	2397-2418	5'-CACGTTCTAATTTTTTTTTAGC-3'	28
	18	2142-2162	5'-AGCCACCGTTCTATTGGCAGC-3'	29
	19	1893-1911	5'-CAACTTCAACTTCTTGATA-3'	30
	20	1142-1165	5'-CCACCACATTCTTTTAAAACTTCC-3'	31
35	21	978-997	5'-GGTAATGGTTCATATGGTGG-3'	32
	22	924-945	5'-GCCATTTGATATAACCATCACC-3'	33
	23	681-702	5'-CTTTAGTTTCGGTTTTAACTTG-3'	34
	24	353-376	5'-CCAAACTGATGAGTAAATTGATGG-3'	35

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* SEQ ID NO:24 and 25 have a one base deletion with respect to the actual sequence shown in FIG. 6. The primers were initally designed from partial sequence infomation and the presence of two "T"s in the actual sequence (positions 2875 and 2932 in FIG. 6) were unknown when the primers were constructed.

3b. Sequence of C. albicans topo I gene

The sequence of the complete open reading frame and surrounding genomic sequences determined is given in SEQ ID NO:1 shown in FIG. 6. Primers used for DNA sequencing are denoted in FIG 6. in bold letters. Potential TATA and CAAT sequences which may be involved in transcriptional initiation are italicized.

3c. Comparative homology of C. albicans topo I to other eukaryotic type 1 topoisomerases

The PILEUP routine of the GCG software package was used to compare the predicted amino acid sequence encoded by the *C. albicans* TOP1 gene with that of topoisomerase I from *S. cerevisiae* (Thrash et al., 1985), *S. pombe* (Uemura et al., 1987) and human cells (D'Arpa et al., 1988). The predicted amino acid sequence determined for *C. albicans* topo I shows 62% overall identity to the predicted amino acid sequence of *S. cerevisiae* topo I and 45% overall identity to the predicted amino acid sequence of the human topo I.

25 3d. Amino acid sequences specific to C. albicans topo 1.

Several regions of amino acids in the topo I sequence of *C. albicans* differ from analogous regions in human topo I. A comparision of the predicted *C. albicans* topo I sequence to the human topo I sequence was performed using the GCG PILEUP routine and is provided in FIG. 7. (According to this program, the symbol "I" indicates identical amino acids between two sequences (i.e. having a match value of 1.0) and the symbols ":" and "." represents similar amino acids between the two sequences (having match values of 0.5 and 0.1 respectively). As seen in FIG. 7, the 5' portion of *C. albicans* TOP1 gene encodes a stretch of charged, hydrophilic amino acids (i.e. from about amino acid position 1 to about position 150) which differ from the amino acids in the 5' amino terminal domain of human topo I. Thus the 5' amino terminal domain of the *C. albicans* enzyme (or nucleotide sequence) may provide a peptide (or nucleic acid) marker specific for *Candida* species.

The predicted amino acid sequence of the C. albicans topoisomerase I also differs

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from the human amino acid sequence in two regions which are associated with camptothecin-resistance in mutants of topoisomerase I from human or S. cerevisiae (Fujimori et al., 1994; Levin et al., 1993), and may contribute to the difference in camptothecin sensitivity seen between topoisomerase I from human and C. albicans (Fostel et al, 1992). Camptothecin has been shown to stabilize the topoisomerase I/DNA cleavage complex, and this stabilization is thought to be the mode of action of this cytotoxic and fungicidal compound. Regions of the topoisomerase protein which are correlated with resistance to camptothecin may be important targets for fungal specific topoisomerase stabilizing agents.

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Two positions associated with camptothecin-resistance in other topoisomerase I genes correspond to (1) the THR residue at position 309 (corresponding to one change observed in a camptothecin-resistant variant of human topoisomerase I reported by Fujimori et al., 1994) and (2) the stretch of amino acids from 728 to 734 which encompasses the active site, and which differs from the homologous human topoisomerase sequence in 2 of the 7 amino acids. Changes in amino acids in this region have also been reported to correlate with camptothecin resistance in S. cerevisiae (Levin et al., 1993)

It should be noted that other segments of the *C. albicans* topo I which do not have homologous partners in the human enzyme may also be important for therapeutic targeting of the *Candida* topoisomerase. There are four regions of this nature, specifically, the amino acids corresponding to positions 573-594, 609-633, 653-677 and 706-712 shown in FIG. 7. Not only are these fungal-specific segments, they are also situated between two segments predicted to be important for catalysis. Thus, their position in the molecule may allow these fungal-specific elements to modulate the enzyme structure or function close to the catalytic center.

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Although it is possible to find amino acid correspondence between the shorter segment in the human gene and the stretch of fungal sequences between 573-712, these correspondences are not strong. Thus, the entire stretch from amino acid 573-712 is structurally different from the shorter stretch in the corresponding position of the human gene. In addition, the particular amino acids which make up this structure differ among C. albicans, S. cerevisiae, and S. pombe.

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The active site of other type I topoisomerases contains a TYR residue which becomes covalently linked to DNA when the catalytic reaction is interrupted (Eng et al., 1989, Lynn et al., 1989). There is a TYR residue at the analogous position in the C. albicans topo I (residue 733 in FIG. 7). The short amino acid sequence surrounding this residue (i.e. SK-X-NY) is conserved in other eukaryotic type I topoisomerases; only the C. albicans topo I gene encodes a MET residue (i.e. SKMNY) rather than the highly conserved ILE or LEU found in other type I topoisomerases (i.e SKINY or SKLNY).

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Amino acids 297-529 span a region of *C. albicans* topo I showing high homology to other eukaryotic topoisomerases. Evidence from other sources suggests that this region contains three domains which in addition to the active site (i.e. the region containing the SK-X-NY motif), are likely to be important to the catalytic function of topo I. (Gupta, M. et al., 1995).

EXAMPLE 4

Expression of C. albicans topo I gene in S. cerevisiae

10 4a. Subcloning of C. albicans topo I gene in an S. cerevisiae expression vector

The C. albicans topo I gene was subcloned into an S. cerevisiae expression vector using standard methods of recombinant DNA technology according to the schematic outline shown in FIG. 9. The C. albicans topo I gene was excised from the lambda clone #26 and moved into the pVT100-U S. cerevisiae expression vector (Thierry et al., 1987), which allows the C. albicans gene to be under the control of the S. cerevisiae alcohol dehydrogenase I (ADH1) promoter. The majority of the C. albicans topoisomerase I gene was excised by cutting the lambda clone #26 with Xba I and Pvu II. This DNA fragment is missing short fragments of the 5' and 3' ends of the open reading frame, so an adapter DNA containing the missing sequences from the 5' and 3' ends of the gene was synthesized. The sequence of the adapter is given in FIG. 8.

The first step in the construction was to insert the adapter DNA into pVT100-U. For this construction, pVT100-U was digested with HinDIII and BamH1 and electrophoresed on an 0.8% low melting point agarose gel (LMPA, GIBCO BRL, Gaithersburg, MD). The agarose slice containing the DNA was melted, added to ligation mix containing the hybridized adapter oligonucleotides (see FIG. 8) with T4 DNA ligase and incubated at 16°C for 24 hours. Competent DH5 α E. coli (GIBCO BRL, Gaithersburg, MD) were then transformed with the reaction mix. Plasmid DNA was isolated as described by Sambrook et al. (1989), and restricted with EcoRI to identify clones containing the adapter DNA. Clone pVT100-UCT7 contained the adapter DNA by this criterion.

The next step in the construction was to insert the *C. albicans* gene in the adapter in pVT100-UCT7. To do this, pVT100-UCT7 was cut with Xba I and Pvu II and electrophoresed on an 0.8% LMPA gel. Lambda clone #26 carrying the *C. albicans* insert was prepared as described by Sambrook *et al.* (1989), and also restricted with Xba I and Pvu II. The lambda DNA was electrophoresed on LMPA, and the agarose containing a 2.3 kb band excised and ligated with T4 DNA ligase at 16° C for 24 hours to the cut pVT100-UCT7 DNA. The ligation mixture was used to transform competent DH5 α *E. coli* (GIBCO

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BRL, Gaithersburg, MD). Plasmid DNA was prepared from several clones as before and digested with Xba I and Pvu II to verify the presence of the 2.3 kb fragment of *C. albicans* DNA. Clone pVT100-UCT7-7, containing this fragment, was selected for further analysis. A culture of *E. coli* DH5α which contains pVT100-UCT7-7 was deposited under the terms of the Budapest Treaty with the Agricultural Research Culture Collection, Peoria, Illinois, on October 11, 1995, and has received the accession number NRRL-B-21504.

4b. Expression of the gene encoding C. albicans topo I in S. cerevisiae

DNA from clone pVT100-UCT7-7 was prepared from a 100 mL culture of E. coli using a Wizard Maxi-prep kit obtained from Promega (Madison, WI) and used to transform S. cerevisiae strains JN362a t1 and JN394 t1 using the method of Schiestl and Gietz (1989). Strains were obtained from Dr. John Nitiss, Children's Hospital, Los Angeles; genotypes are as given below:

JN362a t1: MATa ura3-52, leu2, trp1, his7, ade1, top1::LEU2 ISE2 and

JN394 t1: MATa ura3-52, leu2, trp1, his7, ade1, top1::LEU2 rad52::TRP1 ISE2. The ISE mutation alters cell permeability thereby allowing the strain to be measured for sensitivity to topoisomerase agents such as camptothecin (Nitiss and Wang, 1988). The top1::LEU 2 mutation results in the elimination of functional topo I activity.

Approximately 5 μg of pVT100-UCT7-7 DNA was used for each transformation with 50 μg sonicated denatured salmon sperm DNA (obtained from Dr. Robert Simmer, Abbott Laboratories) added as carrier. Other reagents were obtained from Bio101 (Vista, CA). Cells were selected on SD-ura media (0.67 % w/v Yeast Nitrogen Base from Difco, Detroit, MI; 0.2 % w/v CSM - ura from Bio101, Vista, CA; 2% glucose from Difco, Detroit, MI; 25 mM Hepes, pH 7.2, from Sigma, St. Louis, MO). Ten transformants were selected for further testing, and where determined, showed similar characteristics. Cells carrying pVT100-UCT7-7 grew with similar rates as parent cells carrying pVT100-U (no topo I).

The sensitivity of the transformed cells to camptothecin was determined as one measure of topoisomerase activity. These cells have been shown to be sensitive to camptothecin only when they contain a functional topoisomerase (Nitiss and Wang. 1988). Cells were grown overnight in SD-ura, diluted to a density of 5 x 10⁴ cells/mL as determined by microscopic examination using a hemocytometer, and 200 µL plated in each well of a microtiter plate (Becton Dickinson and Co., Lincoln Park, NJ). Test wells also received 4 µL of different solutions of camptothecin (Sigma, St. Louis, MO) dissolved in DMSO. The minimal inhibitory concentrations for 50% growth inhibition (MIC₅₀) was determined by visual observation after three days of growth at 30°C, and are given in Table 4.

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As shown in Table 4, strains JN362 tl and JN 394 tl, either alone or carrying only the pVT100-U plasmid (i.e. the expression vector lacking a TOP1 gene), were essentially resistant to camptothecin as evidenced by MIC50 values of greater than 100 μ g/mL. In contrast, strain JN394 tl carrying either the *C. albicans* TOP1 gene (in pVT100-UCT7-7), the *S. cerevisiae* TOP1 gene under the control of the ADH1 promoter (in pVT100-UBHT-5) or the human TOP1 gene (in pVT100::HsTOP1) showed MIC50 values of 0.3, 0.3 and 0.005 μ g/mL respectively indicating that these strains were sensitive to camptothecin. Similarly, strain JN362 tl carrying either pVT100-UBHT-5 or PVT100::HsTOP1 showed sensitivity to camptothecin as evidenced by MIC50 values of 10.0 and 0.2 μ g/mL respectively. A transient inhibition of growth was also observed in strain JN 362a tl carrying pVT100-UCT7-7 indicating a partial or lesser degree of sensitivity to camptothecin. For comparison, strain JN394 (having a wild-type level of topo I) was shown to have an MIC50 of 1.56 μ g/mL.

TABLE 4
MIC 50 values (in µg/mL) for S. cerevisiae strains expressing different TOP1 genes.

	No Plasmid	pVT10:0-U	pVT100-UCT7-7	pVT100-UBHT-5	pVT100::HsTOP1
JN 362a tl	>100.0	>100.0	>100.0*	10.0	0.2
JN 394 tl	>100.0	>100.0	0.3	0.3	0.005
JN 394	1.56	-		_	

^{*}transient inhibition of growth was observed.

4c. Characterization of recombinant C. albicans topo I activity

Cells from frozen stocks were streaked on SD-ura with 2% Bacto agar (Difco, Detroit, MI). Individual colonies were used to inoculate 100 mL of SD-ura, grown overnight, then used to inoculate 750 mL cultures at approximately 10.5 cells per mL. After growth for 16 hours, cells were harvested by centrifugation, washed with TGE buffer, and frozen on dry ice. Cells were thawed and lysed by vortexing in the presence of an equal volume of washed .5 mm diameter glass beads (Biospec Products, Inc., Bartlesville, OK) in the presence of TGE buffer containing 1 mM PMSF (phenylmethyl-sulfonyl fluoride), 1 µg/mL aprotinin and leupeptin and 0.1 mM DTT (dithiothreitol), 0.05% v/v Tween 80, all obtained from Sigma, St. Louis, MO.

After approximately 40-45% of the cells were judged to be lysed by microscopic examination, the lysate was recovered and centrifuged at 12,000 rpm in a Sorvall SS34 rotor for 40 minutes at 6°C. The supernatant was recovered, and incubated for 140 minutes at 4°C

with hydroxylapatite resin. The resin was recovered by centrifugation in an IEC refrigerated centrifuge at 4000 rpm using a type 224 rotor for 6 minutes. The resin was washed twice with buffer G plus 0.3 M KPO₄, pH 7.5 (buffer G is 0.1 mM EDTA, 10% glycerol) and collected as before. The topoisomerase activity was eluted from the resin by incubating at room temperature for 10 minutes in the presence of buffer G plus 0.7 M KPO₄ and removing the resin by centrifugation. The eluate was dialyzed overnight at 4°C vs. 50% glycerol, 100 mM KCl, 50 mM Tris pH 7.5, 0.1 mM EDTA, 0.1 mM DTT, 1 mM PMSF and stored at -20°C.

The protein content of each lysate was determined using the method of Bradford (1976) using reagents obtained from Bio-Rad (Hercules, CA), and is given in Table 5. Topoisomerase activity was determined using standard conditions (Fostel *et al.*, 1992). Briefly, dilutions of each lysate were made in a dilution buffer containing bovine serum albumin (BSA; molecular biology grade, obtained from GIBCO BRL, Gaithersberg, MD) in buffer G with 15 mM KPO₄ pH 7.5. One µL of each dilution was added to a 20 µL reaction containing 25 mM Tris pH 7.5; 50 mM KCl; 2.5 mM MgCl₂; 0.5 mM EDTA; BSA at 50 mg/mL; substrate DNA (supercoiled plasmid pBR322 obtained from GIBCO BRL, Gaithersberg, MD) at 6 µg/mL. Reactions were incubated at 30°C for 15 minutes, then stopped by the addition of 3 µL loading buffer (0.67% SDS; 67 mM EDTA; 26.7% sucrose; 0.067% each bromphenol blue and xylene cyanol FF obtained from Sigma, St. Louis, MO), and analyzed by visual inspection after electrophoresis on a 1% agarose gel in TBE buffer (from Mallinckrodt GenAR, Paris, KY). One unit of activity (U)was defined as that needed to relax 50% of the substrate DNA under these conditions. Activity measures are also given in Table 5 below.

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TABLE 5

Strain	Plasmid	Volume (μL)	Protein (μg/μL)	Topoisomerase (U/μL)	Specific Activity (U/µg)
JN 362a t1	pVT100-U	315	6.1	<1	<0.16
JN 362a t1	pVT100-UCT7-7	205	10.7	1000	93
JN 362a t1	pVT100-UBHT-5	225	6.5	300	46

As Table 5 indicates, the presence of the *C. albicans* gene on pVT100-UCT7-7 correlates with topoisomerase activity in the cell.

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We claim:

- 1. An isolated and purified single- or double-stranded polynucleotide selected from the group consisting of
- a) a polynucleotide having a nucleotide sequence comprising the sense sequence of SEQ ID NO:1 from about nucleotide position 345 to about nucleotide position 2807;
- b) a polynucleotide having a nucleotide sequence comprising the sense sequence of SEQ ID NO:1 from about nucleotide position 474 to about nucleotide position 2807;
- c) a polynucleotide having a nucleotide sequence comprising the sense sequence of SEQ ID NO:1 from about nucleotide position 474 to about nucleotide position 923;
- d) a polynucleotide having a nucleotide sequence comprising the sense sequence of SEQ ID NO:1 from about nucleotide position 2199 to about nucleotide position 2624;
- e) a polynucleotide having a nucleotide sequence comprising the sense sequence of SEQ ID NO:1 from about nucleotide position 2655 to about nucleotide position 2807;
- f) a polynucleotide complementary to one of the polynucleotides of (a), (b), (c), (d) and (e);
- g) a polynucleotide that, on expression, encodes a polypeptide encoded by one of the polynucleotide of (a), (b), (c), (d) and (e); and
- h) a polynucleotide that hybridizes, under stringent conditions, to at least one of the polynucleotides of (a), (b), (c), (d), (e), (f) and (g).
- 2. An expression vector comprising an isolated and purified polynucleotide of Claim 1 wherein the polynucleotide is a DNA molecule.
- 3. A host cell transformed with the expression vector of Claim 2.
- 4. The transformed host cell of Claim 3 that is a eukaryotic cell.
- 5. A recombinant C. albicans topoisomerase 1.
- 6. The topoisomerase of Claim 5 having the amino acid sequence selected from the group consisting of SEQ ID NO:2 and SEQ ID NO:3.

- 7. An isolated and purified polypeptide of fewer than about 821 amino acid residues having an amino acid sequence comprising at least one of
- a) the sequence from residue position 1 to residue position 150 of SEQ ID NO:3;
- b) the sequence from residue position 573 to residue position 717 of SEQ ID NO:3; and
- c) the sequence from residue position 728 to residue position 778 of SEQ ID NO:3.
- 8. A method for identifying antifungal activity in a test compound comprising the steps of:
 - (a) providing a first living cell and a second living cell;
- (b) altering said first living cell to produce a modified living cell, wherein said modified living cell has an increased level of topoisomerase I activity relative to said second living cell;
- (c) exposing said modified living cell and said second living cell to said test compound under conditions suitable for cell growth; and
- (d) comparing the extent of cell mortality or attentuated cell growth of said modified living cell to said second living cell.
- 9. The method of Claim 8 wherein said step of altering said first living cell is achieved by introducing into said first living cell an expression vector or plasmid comprising a polynucleotide having a nucleotide sequence comprising the sequence of SEQ ID NO:1 from about nucleotide position 474 to about nucleotide position 2807.
- 10. A method for identifying antifungal activity in a compound comprising the steps of:
 - (a) providing a first living cell and a second living cell;
- (b) altering said first living cell to produce a modified living cell, wherein said modified living cell has a decreased level of topoisomerase I activity relative to said second living cell;
- (c) exposing said modified living cell and said second living cell to said test compound under conditions suitable for cell growth; and
- (d) comparing the extent of cell mortality or attentuated cell growth of said modified living cell to said second living cell.
- 11. The method of Claim 10 wherein said step of altering is achieved by disrupting a

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coding region of a chromosomal topoisomerase I gene in said first living cell by introducing into said first cell a polynucleotide comprising a selectable marker gene cloned within a polynucleotide having a nucleotide sequence which comprises a sequence selected from the group consisting of

- (a) SEQ ID NO:1 from nucleotide position 345 to nucleotide position 2807;
- (b) SEQ ID NO:1 from nucleotide position 474 to nucleotide position 2807; and (c) a fragment of (a) or (b).
- 12. The method of Claims 8 or 10 wherein said first living cell and said second living cell are both derived from the same strain of S. cerevisiae or C. albicans.

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FIG 1

Hu	FRTYNASITLQ(QLKELTAPE	ENIPAKI	LS	YNR	ANRA	VAILCNHQ	₹
	111111	1	l.	I	11	111	*****	l
Sc	FRTYNASKTMQ1							
	1111111	l	1	1	11	Ш	11111111	İ
Sp	FRTYNASYTMA	EELKKMPKNI	TLAD.KI	LF	YNR	ANRI	rvailcnhq1	R
_	1111111 1	l	1	ı	11	111	111111111	1
Dm	FRTYNASKTLO	SOLDLLTDPS	SATVPEKL	L	YNE	ANRA	VAILCNHQ	R

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FIG. 2

CGCAGAGAAAATATTGAAGTACAACGCAGCAAATAGAACTGTAGCCATCCTATGTAACCATCAAAGGA TYTYCCGTACATATAATGCTTCCAAAACAATGCAAGATCAACTGGATCTAATTCCAAATAAAGGATCTGT **CGTACATATAATGCCTCGAAAACCATGCAAGATCAAATTGATATAATTGAAAATGAAGGTACAGT GCCGGAAAAAGTGGCTAAATTCAATGCTGCCAATAGAACGGTGGCTATTCTTTGCAATCACC** - - - - - -Sc: Sc: S C B Ca:

FIG.

RITYNASITLQ QQL....DEL TNSDDNVPAK I.....LS YNRANRAVAI LCNHQRA KVF RTYNASYTMA EEL...KKM .PKNLTLADK I.....LF YNRANRTVAI LCNHQRS KVF RTYNASVTFQ GLLEQTEE.W LKSRPNAAER EINQTNLRLA YNEANRQVAI LCNHQKT KVF RTYNASITLQ QQL....KEL TAPDENIPAK I.....LS YNRANRAVAI LCNHQRA KVF RTYNASITLQ QQL....KEL TAPDENVPAK I.....LS YNRANRAVAI LCNHQRA RTYNASKTLQ SQL....DLL TDPSATVPEK L.....LA YNRANRAVAI LCNQRSV RTYNASITLD EML....SQE T.KDGDVTQK I......VV YQKANKEVAI ICNHQRT RTYNASKTMQ DQL...D.L IPNKGSVAEK I.....LK YNAANRTVAI LCNHQRT ... RTYNASKTMQ DQI...D.I IENEGTVAEK V.....AK FNAANRTVAI LCNH. KVF KVF KVF KVF E 톳 Ę H SC Sp X Ç

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FIG. 4

Hu IFRKNFFKDW
Mm IFWKNFFKDW
Dm VFNNNFFKDF
SC VFQKNFFNDF

FIG. 5

1 AAGAATTTTT TCAATGATTT TTTGGAAGTT TTAAAAGAAT GTGGTGGTTG TGGTGTTGAA ATTAAAAAAT TTGAAAAATT AGATTTTAGT AAAATGTATG CTCATTTTGA AAAATTACGT GAAGAGAAAA AAGCCATGAG TAGAGAAGAA AAGAAAGAA TCAAAGAAGA AAAAGAAAAA GAAGAAGAAC CTTATAGGAC TTGTTATCTT AATGGTAGAA AAGAATTAGT GGGGAATTTC CGTATTGAAC CTCCAGGTTT ATTCCGTGGT CGTGGTGCAC ATCCTAAAAC TGGGAAATTG AAACGTCGAG TAGTGCCGGA ACAAGTGACT TTGAATTTAG GTAAAGATGC 301 TAAAATACCT GAACCACCAG CAGGTCATCA ATGGGGGGAA ATTAGACATG ATAATGAAGT CACTTGGTTG GCCATGTGGA AAGAAAATAT TTCTGATTCG 401 TTGAAATATG TTAGATTTGC TAATAATTCA TCAGTTAAAG GACAATCTGA 451 TTTCAAAAA TTTGAAACGG CGAGAAAATT AAAGAGAGAT CACGTTGATT 501 CTATTAGAAA AGATTATACC AAAATGTTAA AATCAGAGAA AATGCAAGAT 551 AGACAAATGG CCACGGCTAT GTATCTTATT GATGTTTTTG CATTGAGGGC 601 TGGTGGTGAA AAAGGTGAGG ATGAAGCCGA TACCGTTGGT TGTTGTTCAT 651 TACGATATGA ACATGTAACT TTAAAACCAC CCAACAAGGT TATTTTCGAT 701 CTTTTGGGTA AAGATTCAAT TAGATTTTAT CAAGAAGTTG AAGTTGATAA 751 ACAAGTTTTC AAAAATCTAC GAATTTTCAA AAAATCTCCT AAACAACCTG 801 851 GTGATGATTT ATTTGATCGT ATAAACCCTT CATTAGTCAA TCGACAATTA 901 CAAAATTATA TGAAAGGATT AACAGCAAAA GTTTTCCGTA CATATAATGC 951 CTCGAAAACC ATGCAAGATC AAATTGATAT AATTGAAAAT GAAGGTACAG 1001 TGGCGGAAAA AGTGGCTAAA TTCAATGCTG CCAATAGAAC GGTG

SEQ ID NO:2

FIG. 6(a)

09	120	180	240	300	360	
TITITIGGTGATGGTTTTAGGCGCGCGATTAATCATTTTACTATTGAGAATGTATGCTCC 	Cacatictatiacacctcatctttcatctttcatcttttcatctntcaatcactaaatalaa 	CCTTGCGACCTTCACAAATTTTTTTTTTTTGACAAGCAATCCAAAATTACAATTTTCATTT 	CATTTCTTTTATATATAAAAGTTTTTCACCATTAATTTCACCACACACTCTCATTAGCAAT	TGGGCAAAATAGAAAGTAATTT <i>TATAA</i> CTTATAACCAAAAACAATTCAAGAACAATAT 	Cattattattaaatttatcacggaattigititaaatcaagtaagaacaattcca <i>tcaat</i> 	
~	61	121	181	241	301	
EQ ID NO:1						
I Oas						

FIG. 6(b)

361	TTACTCATCAGTTTGGTTGTAATAATAAAACAGATTATTTTTTTT	420
	THOFGCNNKNRLFFLSSPPR-	
	GAGTATTCCGTTATTTAAGTCCATTATTTGTTCGTTCATATAGCATAATTCCTATGAATT 421++++++	480
	VFRYLSPLFVRSYSIIPMNS-VFRYLSPLFVRSYSIIPMNS-	
481	CATCAGACGAAGAAGACATTGCCTTGTCTAGACTTGCTAAAAAATCATCCTCGATCACTT	540
	S D E E D I A L S R L A K K S S S I T S -	
541	CAGCTTCCACTTATGAAGACGATGAAGATGATGATATCCCTTTGGCTAAAAATCCAGGA	600
	ASTYEDDEDDDIPLAKKRR.	
601	AAAAGAGGGTTGAATCTGATTATGAAGAGATGAAGACGAAGTCCCATTGAAAAGTTGT 	999
	KRVESDYEEDEDEVPLKKLS-	

FIG. 6(c

661	CTAATGGTAGAGCAAAAAACAAGTTAAAACCGAAACTAAAGTTAAAAAGGAACCTAAAA ++++++	720
	NGRAKQVKTETKVKKEPKS-	
121	GTGCCAATAAATCCAAATCTACATCTAAAAAGGACACCAAAGTTAAGAAAGA	780
12)		
	ANKSTSKKDTKVKKEKTT-	
1	CAGTCAAGAAGGAATCTAAAGCCACACAAGCACTAAAGTGAAAGAAGAATCCAAAACTCAAT	048
781	GTCAGTTCTTCCTTAGATTTCGGTGTTTCGTGATTTCACTTTCTTT	5
	V K K E S K A T S T K V K E E S K T Q S -	
	CAGATTCACAAGCATCGGTTAAATCTGAAACTCCTGAAGAAGATCAAGGGTACAAATGGT	C
841	GICTAAGICITCGTAGCCAATTTAGACTTTGAGGACTTCTTCTAGTTCCCATGTTTACCA	
	DSQASVKSETPEEDQGYKWW-	
	GCGAAGTGAATCAAGAAGAAGGTGATGGTTATATCAAATGGCAAACACTAGAACATA	•
901	ccttcacttagttcttcttcttccactaccaatatagtttaccgtttgtattat	966
	H N N O E E E G D G Y I K W Q T L E H N -	

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FA 1020		i Z	AA		ı E		AA 1140		AG 1200		- Q		-+ 1260 AT	۳.
CTITICCACCATATGAACCATTACCATCTCATGTCAAATTATATATA	AAT	HVKLYYN	CAGITAATITTACCTCCAGAAGCAGAAGTIGCCGGATITTATGGAGCAA	TCGJ	æ	TII	CTAAAAA	A K N P V F Q K N F F N D F L	taaaagaatgiggiggtigtggttigaaatttaaaaatttgaaaatttag	TA	ECGGCGVEIKKFEKL	GAG	CTC	လ
ATA	TAT	>-	TGG	ACC.	O	TGA TGA	ACT	۵	AAA	111	×	CAT	GTA	Σ
ATT	TAA	ı	TTA	AAT	>	%	CTT	z	TC	ACT	凹	NA GO	÷ L	æ
CA.	GTT	×	SATT	T.	A E E V A G F Y G	E	PACGATTTTTGGGTCAAAAGGTTTTTCTTAAAAAAGTTA	Ĺ	AATT	T.	ţĿ	A.		×
ATG:	FACA	>	8	200	O	ATT	T.	ţri	AAA	11.1	×	AGA	TCT	
CTC.	3861	=	TTC	AACC	A	AGA	TCT	Z	TTA	AAT	×	AAG	TIC.	ы
CAT	GTA	Sdldaydda	Z AG	TIC	>	A.A.	1	×	AAA :	11.	H	GTG	CAC	ш
TAC	ATG	D.	AAG	TTC	េ	TCC	AGG	O ¹	TTG	MAC	Щ	TAC	ATG	K L R
CAT	GTA	<u>, 1</u>	CAG	GTC	М	I.I.	* A	<u>[14</u>	50.10	CAC	>	AA1	Į.	
AAC	TTC	a.	AAG	Tric	а M	CAG	GTC	>	IGTG	ACAC	0	18		ы ж
ATC	TAC	ш	CAG	GTC) ∑) E	7	SGT	Z S	()	FI	A A	Ĺ.,
CAT	i i	۸.	CTC	3GAC	<u>ω</u>	₹		~ ×	SGT	CCA	C)	CAT	GTA	*
CAC	SCTC	۵.	ITAC	AATC	بر د	71.05 71.05	CGA	~ «	TGT	ACA	Ü	g	CGA	
CCAC	SGT	Δ.	AAT	TTA	V N L P P	CAT	GTA	×	GAA	CTT	ы	TAT	ATA	M Y A H
TTT	A.A.	E4	GIT	CAA	>	GAT	CTA	H Q	AAA	TTI	×	LATG	TAC	Σ
ATG	TAC		CC.	GGT	Δ.	ACT	TGA	E	ATT.	183	ដ	2	ŧĘ	×
CTT	80	∑ >	Y. A.A.	Tati	×	GAA	ÇİJ	ω	CTI	ฎี	>	[3]	ATC.	S
ACGCIGITATGITTCCACCACCATATGAACCATTACCATCTCATGICAAATTATATATA	TCCCACAATACAAAGGTCGTGTATACTTGGTAATGGTAGAGTACAGTTTAATATATAT	ပ	ACAATAAACCAGTTAATTTACCTCCAGAAGCAGAAGAAGTTGCCGGATTTTATGGAGCAA	TCTTATTTCGTCAATTAAATCGAGGTCTTCGTCTTCTTCAACGGCCTAAAATACCTCGTT	Z	TGTTAGAAACTGATCATGCTAAAAACCCAGTTTTTCCAAAAGAATTTTTTCAATGATTTTT	ACAATCTTTGACTAGTACGATTTTTGGGTCAAAAGGTTTTTCTTAAAAAAGTTTACTAAAA	٦	TGGAAGTTTTAAAAGAATGTGGTTGTTGTGGTTTGAAATTTTGAAAATTTTGAAAATTTAG	ACCTTCAAAATTTTCTTACACCCACCACACCACTTTTAATTTTTT	ជ	attttagtaaaatgtatgctcatttttgaaaattacgtgaagagaaaaaagccatgagta	TAAAATCAITTTACATACGAGTAAAACTTTTTAATGCACTTCTCTTTTTTGGTACTCAT	Ĺų
130	186			1701		9	1001		-	4 P 4			1201	

FTG. 6(6

1320	1380	1440	1500	1560
GAGAAGAAAGAAAGAAAAAAAAAAAAAAAAAAAAAAAA	GTTATCTTAATGGTAGAAATTAGTGGGGAATTTCCGTATTGAACCTCCAGGTTTAT	TCCGTGGTCGTGCTCCTAAAACTGGGAAATTGAAACGTCGAGTAGTGCTGGAAC+++++++	AAGTGACTTTGAATTTAGGTAAAGATGCTAAAATACCTGAACCACCAGCAGGTCATCAAT+++++ TTCACTGAAACTTAAATTCTACGATTTTTATGGACTTGGTGGTCGTCGAGTAGTTA V T L N L G K D A K I P E P P A G H Q W -	GGGGGGAAATTAGACATGATAATGAAGTCACTTGGTTGGCCATGTGGAAAGAAA
1261	1321	1381	1441	1501

FIG. 6(f.)

ACAATCTGATT ++ 1620 TGTTAGACTAA	Q S D F - TAGAAAGATT ++ 1680 ATCTTTTCTAA	CGCTATGTATC -++ 1740 CCGATACATAG A M Y L -	AAGCCGATACCG -++ 1800 TTCGGCTATGGC A D T V -	ACAAGGTTATTT 1860 -++ 1860 TGTTCCAATAAA K V I F -
CTGATTCGTTGAAATATGTTAGATTTGCTAATAATTCATCAGTTAAAGGACAATCTGATT 	TCAAAAAATTTGAAACGCCGAGAAAATTAAGAGATCACGTTGATTCTATTAGAAAAGATT AGTTTTTAAAACTTTGCCGCTCTTTTAATTCTCTATTGCAACTAAGATAATCTTAAAAATTCTTAAAAATTCTTAAAAAAAA	ATACCAAAATGTTAAAATCAGAAAATGCAAGATAGACAAATGGCCACGGCTATGTATC ATACCAAAATGTTAAAATCAGAAAATGCAAGATAGACAAATGGCCACGGCTATGTATC TATGGTTTTACAATTTTAGTCTTTTTACGTTCTATCTGTTTACCGGTGCCGATACATAG	THATTGATGTTTTTGCATTGAGGCTGGTGGTGAAAAGGTGAGGATGAAGCCGATACCG AATAACTACAAAAACGTAACTCCCGACCACCACTTTTTTCCACTCCTACTTCGGCTATGGC IDVFALR GGEKGEVGTAACTCCCGACACACTTTTTTTCCACTCCTACTTCGGCTATGGC	TIGGINGINGTICATTACGATATGAACATGTAACTITAAAACCACCCAACAAGGTTATIT
	1621	1681	1741	1801

 $(\hat{\mathfrak{g}}) = (\hat{\mathfrak{g}} - \hat{\mathfrak{g}}) \cdot (\hat{\mathfrak{g}} - \hat{\mathfrak{g}})$

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1920	1980	2040	2100	2160
TCGATCTTTTGGGTAAAGATTCAATTAGATTTTATCAAGAAGTTGAAGATAAACAAG	TTTTCAAAATCTACGAATTTTCAAAAATCTCCTAAACAACCTGGTGATGATTTATTT	ATCGTATAAACCCTTCATTAGTCAATTACAAATTATATGAAAGGATTAACAG TAGCATATTTGGGAAGTAATCATTAGCTGTTAATGTTTTAATATACTTTCCTAATTGTC R I N P S L V N R Q L Q N Y M K G L T A -	CAAAAGTTTTC + GTTTTCAAAAG K V F	AAAATGAAGG
1861	1921	1981	2041	2101

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FIG. 6(h)

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			2161	CTATTTTAIGHAICACCIACCGIACCGIACCGIACCGIACCACTATCACAAGTCTTT GATAAAATACATTAGTGGTGGTGCATGCCAGTCATTTTTGGGTACCACTATCACAAGTCTTT T	2220
			2221	TGACAAATTGAAI	2280
			2281	TAGAACCC	2340
<u>ப</u>	Δ.	×	L 234]	K K K D S K Y F E E I D D L I - TCAAAGAAGATATTGAACATATTCATCATACTATAATTAAAAGACAACGAGAACAAGCTA TCAAAGAAGATATTGAACATATTCATCATAATTAAATTTTCTGTTGCTCTTGTTCGAT AGTTTCTTCTATAACTTGTATAAGTAGTATGATATTTAATTTTTCTGTTGCTCTTGTTCGAT	2400
			2401	AAAAAAATTAGAACGTGATAATGAAAATTGAAACTTGAAGGTAAACCATTATTAACTG +++++ TITITITAATCTTGACTATTACTTTTTAACTTTGAACTTCCATTTGGTAATAATTGAC K K L E R D N E K L K L E G K P L L T E -	2460

e 0 X e

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FIG. 6(j

2461	AATCAGATATAAAATTAGATAAAATTGATGAATTAGAAAAGAATATCAAAAAG+++++ TTAGTCTATATTTTCTATTTTAACTACTTAATCTTTTTTCTTATAGTTTTTC C D I K D K L D K I D E L E K E Y Q K E -	2520
2521	GAAAACT +- CTTTTGA K T	2580
2581	TGAAACT	2640
2641	ATAATTCTGAAGTTTCTTTAGGAACTTCAAAAATGAATTATATTGATCCAAGATTAATTG	2700
2701	TRATCTTTTCTAAAAATTTGATGTTCCTATTGAAAATTATTTACCAAAACTTTAAGAG+++++ AATACAAAAGATTTTTTAACAAGGATAACTTTTTAATAATAATGGTTTTGAAATTCTC M F S K K F D V P I E K L F T K T L R E -	2760

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2761	AAAAGTICATTIIGGGCTATIGAATCAGCTGATGAAAATTGGAGATTCTAAAATTAGGGGT	2820
	KFIWAIESADENWRF	
2821	TTGTTTCTTAGCTTATTATTATACTATATGCTGTAGAGTAAATTTTGTACCTTGTAA 	2880
2881	TATATATCCACTITATITICAACATAAAAAATAGATTGATACTGCAGTGTGAAAAGGAAT	2940
2941	ATGCACACCAAGCAAGTATATTTTAGATAAGGATTGGTGTTTTGATATTGGAAGGGC	3000
3001	GAAAGANGAAGGGGGGTATCACATAGACACGTACAATCAAGAAATTGAAATTTTTCCGA 	3060
3061	ATTCCGTTAAATAAAATTTTTACGCGCCTCAAATTTTTCTTCTGTTTTTTGCAAC	3120
3121	TTCTGCAGATAGAGCAACAGTTCCCCAATATCTCTATTGGAAAGATATCACAATCTCAATA	3180
3181	GAGTAATCGTGGGAGAGTCTAATATATACGTATATACTCCTTCA 	

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FIG. 7 (a)

	* * * * *	
2	NSSDEEDIALSRLAKKSSSITSASTYEDDEDDDIPLAKKSRKK	44
55	:	104
45	RVESDYEEDEDEVPLKKLSNGRAKKQVKTETKVKKEPKSANKS	87
105	.:: : . :.: .:: .:: . EKENGFSSPPQIKDEPEDDGYFVPPKEDIKPLKRPRDEDDVDYKP	149
88	KSTSKKDT. KVKKEKTTVKKESKATSTKVKEESKTQSDSQASVKSETPEE	136
150	KKIKTEDTKKEKKRKLEEEEDGKLKKPKNKDKDKKVPEPDNKKKKPKKEE	199
177	DQGYKWWEVNQEEEGDGYIKWQTLEHNGVMFPPPYEPLPSHVKLYYNNKP	186
1,	: : : . -:::: .:: : : :. .	200
200	EQKWKWWEEERYPEGIKWKFLEHKGPVFAPPYEPLPENVKFYYDGKV	246
	+ + + +	
187	VNLPPEAEEVAGFYGAMLETDHAKNPVFQKNFFNDFLEVLKECGGCGVEI	236
	:.1. .	
247	MKLSPKAEEVATFFAKMLDHEYTTKEIFRKNFFKDWRKEMTNEEKNII	294
	+ + + + +	
	KKFEKLDFSKMYAHFEKLREEKKAMSREEKKRIKEEKEKEEEPYRTCYLN	286
	:.	
295	TNLSKCDFTQMSQYFKAQTEARKQMSKEEKLKIKEENEKLLKEYGFCIMD	344
	+ + + + + +	
287	GRKELVGNFRIEPPGLFRGRGAHPKTGKLKRRVVSEQVTLNLGKDAKIPE	336
	.:14 ::11:111111111111111111111::.1::.:1 :1111:1.	
345	NHKERIANFKIEPPGLFRGRGNHPKMGMLKRRIMPEDIIINCSKDAKVPS	394
	+ + + + +	
337	PPAGHQWGEIRHDNEVTWLAMWKENISDSLKYVRFANNSSVKGQSDFKKF	386
	11:11.1 1:1111.1111. 1.111 :1:11:.:1.:11:.1:.1:	
395	PPPGHKWKEVRHDNKVTWLVSWTENIQGSIKYIMLNPSSRIKGEKDWQKY	888
	+ + + + +	
887	ETARKLRDHVDS1RKDYTKMLKSEKMQDRQMATAMYLIDVFALRAGGEKG	436
145	ETARRLKKCVDKIRNQYREDWKSKEMKVRQRAVALYFIDKLALRAGNEKE	494
	+ + + + +	
137	EDE.ADTVGCCSLRYEHVTLKPPNKVIFDLLGKDSIRFYQEVEVD	480
	1:1 111111111 11:.1.1.	
195	EGETADTVGCCSLRVEHINLHPELDGQEYVVEFDFLGKDSIRYYNKVPVE	>44
	+ + + + +	
81	KQVFKNLRIFKKSPKQPGDDLFDRINPSLVNRQLQNYMKGLTAKVFRTYN	530
	1.11111.:1 111:111111:1.:::1::11:.1.11111111	F 4 4
45	KRVFKNLQLFMEN.KQPEDDLFDRLNTGILNKHLQDLMEGLTAKVFRTYN	273

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FIG.7 (b)

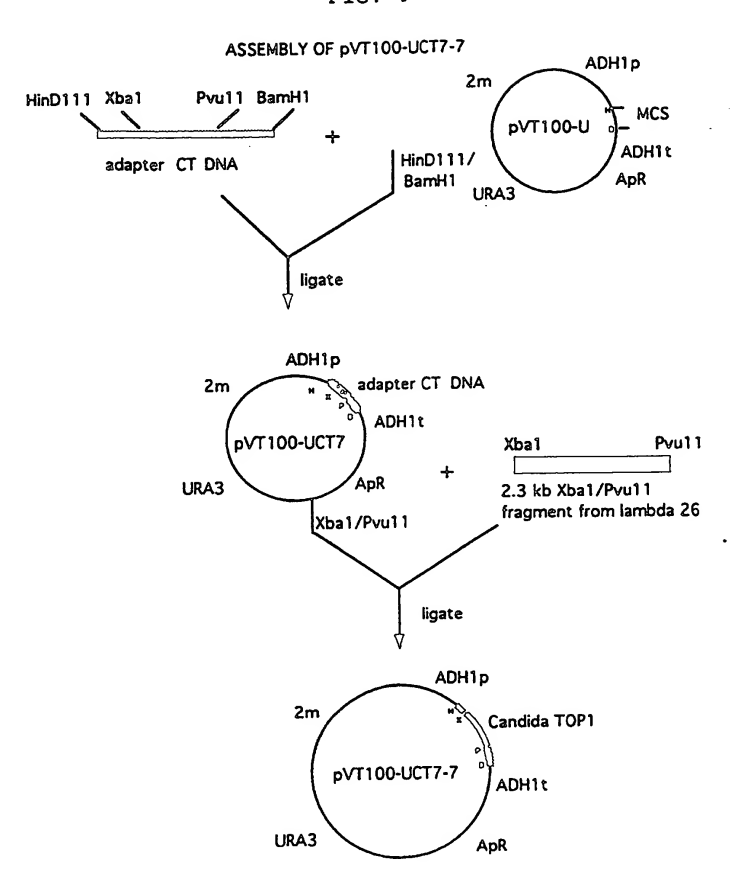
	* * * *	
531	ASKTMQDQI.DIIENEGTVAEKVAKFNAANRTVAILCNHQRTVSKTHGDS	579
	11 1:1:1: :: ::.:. :: .:	
594	ASITLQQQLKELTAPDENIPAKILSYNRANRAVAILCNHQRAPPKT	639
	+ + + + + + + + + + + + + + + + + + +	620
580	VQRINDKLKKFMWQKIRLKKMILQLEPKLKKKDSKYFEEIDDLIKEDIEH	023
	:.1 ::.1:.1	C C /
640	FEKSMMNLQTKIDAK	654
	IHHTIIKRQREQAKKKLERDNEKLKLEGKPLLTESDIKDKLDKIDELEKE	679
630	.: ::. :	•
c = =	KEQLADARRDLKSAKADAKV	674
033	· · · · · · · · · · · · · · · · · · ·	
	* * * * *	
680	YQKELKTGKPIVTKNATVEKLKQQIETLENRILNVSIQLKDKEDNSEVSL	729
	1. 1. 1. 1	
675	.MKDAKTKKVVESKKKAVQRLEEQLMKLEVQATDREENKQIAL	716
•		
	+ + + + +	
730	GTSKMNYIDPRLIVMFSKKFDVPIEKLFTKTLREKFIWAIESADENWRF 7	78
	1111:11:111:1 :: 1 :: 11:: 1111:: . 1 1111 111: 111:: 1	
717	GTSKLNYLDPRITVAWCKKWGVPIEKIYNKTQREKFAWAIDMADEDYEF 7	65

FIG.

5 ' agcttatgaattcatcagacgaagagacattgccttgt<u>ctag</u>aa*tt<u>cag</u>ctg*atgaacattggagattctaaaattag

3 ' ATCTTAAGTAGTCTGCTTCTTCTGTAACGGAACA<u>GATC</u>TTAA<u>GTCGAC</u>TACTTGTAACCTCTAAGATTTTAATCCTAG

FIG. 9



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(57) Abstract

The present invention provides an isolated and purified polynucleotide that encodes Candida albicans type I topoisomerase. Methods of making recombinant C. albicans topoisomerase I using those polynucleotides and host cells transformed with those polynucleotides are also provided. The present invention also provides a method for identifying compounds which inhibit the growth of fungal cells using the polynucleotide.

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INTERNATIONAL SEARCH REPORT

Interr nal Application No PCT/US 96/17291

A. CLASS IPC 6	C12N15/61 C12N15/81 C12N9 C12Q1/533	9/90 C12N1/19	C12Q1/02	
According	to International Patent Classification (IPC) or to both national	classification and IPC		
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	ation searched other than minimum documentation to the extent			
	data base consulted during the international search (name of da	ita base and, where practical, search term	ıs used)	
	MENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of	the relevant passages	Relevant to claim No.	
Υ	CHEMICAL ABSTRACTS, vol. 117, 23 November 1992 Columbus, Ohio, US;	no. 21,	1	
	abstract no. 207692, FOSTEL, JENNIFER M. ET AL: "Characterization of DNA topoi			
	from Candida albicans as a tar discovery" XP002034941	get for drug		
	see abstract & ANTIMICROB. AGENTS CHEMOTHER 36(10), 2131-8 CODEN: AMACCQ;I 0066-4804,	l. (1992), SSN:		
γ	WO 89 09222 A (UNIV JOHNS HOPK October 1989 see claims	(INS) 5	1	
l	See Clathis			
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* Special cat	tegories of cited documents:	"T" later document published after	the international filing date	
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other n	means ent published prior to the international filing date but han the priority date claimed	ments, such combination being in the art. "&" document member of the same	obvious to a person skilled	
	actual completion of the international search	Date of mailing of the internation		
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INTERNATIONAL SEARCH REPORT

International Application No
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